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(54) Title: AN ANTISENSE OLIGONUCLEOTIDE PREPARATION METHOD

#### (57) Abstract

A method for the preparation of an antisense oligonucleotide or derivative thereof comprising the steps of: selecting a target nucleic acid, if necessary elucidating its sequence; generating the antisense oligonucleotide with the proviso that: the oligonucleotide comprises at least 8 residues; the oligonucleotide comprises at maximum twelve elements, which are capable of forming three hydrogen bonds each to cytosine bases; the oligonucleotide does not contain four or more consecutive elements, capable of forming three hydrogen bonds each with four consecutive cytosine bases (CCCC) within the target molecule or alternatively four or more consecutive elements of GGG; the oligonucleotide does also not contain 2 or more series of three consecutive elements, capable of forming three hydrogen bonds each with three consecutive cytosine bases (CCC) within the target molecule, or alternatively 2 or more series of three consecutive elements of GGG; and the ratio between residues forming two hydrogen bonds per residue (2H-bond-R) with the target molecule and those residues forming three hydrogen bonds per residue (3H-bond-R) with the target molecule, is ruled by the following specifications: 3H-bond-R/3H-bond-R/4H-

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# An antisense oligonucleotide preparation method

The present invention is related to a method for the preparation of antisense oligonucleotides and to an oligonucleotide or functional or structural analogs or effective derivatives thereof, forming hydrogen bonds with deoxyribonucleic acids (DNA) and/or ribonucleic acids (RNA) or derivatives thereof including, but not limited to the formation of hydrogen bonds with the bases adenine (A), cytosine (C), guanine (G), uracil (U) or thymidine contained in such molecules or forming hydrogen bonds with residues of a particular protein, such a molecule being capable of altering the expression structure or function, of a gene, an RNA molecule or a protein or altering the level of activity of a gene, an RNA molecule or a protein. Furthermore, the present invention is related to such nucleic acid or functional or structural analogs or effective derivatives thereof, coupled or mixed with folic acid, hormones, steroid hormones such as oestrogen, progesterone, corticosteroids, mineralocorticoids, androgens, peptides, proteoglycans, phospholipids, glycolipids and derivatives therefrom.

Furthermore, the invention is related to the use of said nucleic acids or functional or structural analogs or effec-

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tive derivatives thereof, for analyzing the functional properties of a particular gene, RNA, or protein by altering its activity, structure, function or altering its expression levels.

Furthermore, the invention is related to antisense nucleic acids, capable of modulating the expression or functional activity of proteins which regulate cell growth leading to augmentation, inhibition or modulation of cell growth or cell proliferation and/or the expansion of primary cells or stem cells, e.g. in culture or in the living organism.

Furthermore, the invention is related to a pharmaceutical composition comprising said nucleic acids or functional or structural analogs or effective derivatives thereof, hybridizing with an area of the messenger RNA (mRNA) or the DNA of a target gene or binding to a particular protein as well as the use of said nucleic acids, structural analogs and derivatives thereof for the manufacturing of a pharmaceutical composition for the treatment of diseases where the alteration of the structure function, activity or expression of a particular target gene, a particular target RNA or a particular target proteins activity leads to a therapeutic benefit related to the effect of the nucleic acid or derivative thereof.

Modulation of the expression of genes, RNA molecules or proteins or of their activity levels with nucleic acids or functional or structural analogs or effective derivatives thereof is a powerful means to study the function of the respective molecules. For example modulation, e. g. knockdown or increase of the expression of a particular protein can lead to the identification of its physiological as well as its pathophysiological roles in cultured cells as well as in living organisms in vivo.

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Furthermore, the aberrant expression or overexpression of genes, RNA molecules or proteins, the expression of foreign DNA, RNA or proteins e. g. derived from infectious organisms or the expression of mutated DNA, RNA and proteins is found in a variety of diseases. Downregulation of the expression or the activity of such DNA, RNA and/or proteins can lead to an inhibition of or to the reversal of pathological processes in which the expression of a particular DNA, RNA and/or protein plays a role. However, nucleic acids or derivatives thereof used for downregulation of DNA, RNA and/or protein expression are often ineffective and/or toxic to the cells or the organisms treated with such molecules.

An object of the present invention is to provide a method for designing and preparation of oligonucleotides or derivatives thereof which avoid the drawbacks of prior art, and give a reliable method for preparation of oligonucleotides having increased effectivity and/or reduced toxicity and/or reduced non-selective effects.

The object is attained by a method having the features of claims 1. Preferred embodiments of the method of the invention are those according to claims 2 to 7.

The method of the invention comprises the steps

- of selecting a target nucleic acid, if necessary elucidating its sequence
- generating the antisense oligonucleotide with the proviso that
  - the oligonucleotide comprises at least 8 residues,
  - the oligonucleotide comprises at maximum twelve elements, which are capable of forming three hydrogen bonds each to cytosine bases,

- the oligonucleotide does not contain four or more consecutive elements, capable of forming three hydrogen bonds each with four consecutive cytosine bases (CCCC) within the target molecule or alternatively four or more consecutive elements of GGGG,
- the oligonucleotide does also not contain 2 or more series of three consecutive elements, capable of forming three hydrogen bonds each with three consecutive cytosine bases (CCC) within the target molecule, or alternatively 2 or more series of three consecutive elements of GGG, and
- the ratio between residues forming two hydrogen bonds per residue (2H-bond-R) with the target molecule and those residues forming three hydrogen bonds per residue (3H-bond-R) with the target molecule, is ruled by the following specifications:

3H-bond-R

 $\geq 0.29$ 

3H-bond-R + 2H-bond-R

and synthesizing the oligonucleotide thus generated in a per se known manner.

The generated antisense oligonucleotide comprises at least 8 residues in order to have sufficient interaction with the target molecule and has preferably up to 30, more preferably up to 24 or most preferred upt to 18 residues. Shorther chain length are preferred over longer ones to increase specifity and/or reduce non-specific effects.

The oligonucleotide comprises at maximum 12 elements which are capable of forming 3 hydrogen bonds each to cytosine bases. In case of generating an oligonucleotide an element is represented by a residue, thus a nucleotide of the oligo-

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nucleotide. In cases of generating a derivative an element is considered as a part of the molecule capable of forming hydrogen bonds. It is preferred that the oligonucleotide comprises at maximum 10 and more preferred at maximum 8 elements which are capable of forming 3 hydrogen bonds each to cytosine bases.

The generated antisense oligonucleotide preferably does not contain 4 or more consecutive guanine bases and does also not contain 2 or more series of 3 consecutive guanine bases.

Preferably, the ratio between residues forming 2 hydrogen bonds per residue (2H-bond-R) with their target molecule and those residues forming 3 hydrogen bonds per residue (3H-bond-R):

3H-bond-R

3H-bond-R + 2H-bond-R

is in the range of greater than 0.33 and smaller than 0.86, more preferably smaller than 0.79 and still more preferred smaller than 0.72.

In one embodiment the oligonucleotides generated by the method of the invention are modified for higher nuclease resistance than naturally occurring nucleotides. Methods for synthezing oligonucleotides and derivatives thereof are known in the art, see for exammple "Oligonucleotides and Analogues", F. Eckstein (Ed.), 1991, IRL Press Oxford or "Protocols for Oligonucleotides and Analogs, Synthesis and Properties", Sudhir Agrawal (Ed.), 1993, Humana Press, Totowa, New Jersey.

Oligonucleotides of the invention may also contain RNA and DNA residues within their chains.

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linkages preferably consist of methylphosphonate linkages or phosphodiester linkages.

The chemical structures of antisense oligodeoxy-ribonucleotides are given in figure 1.

The chemical structures of antisense oligo-ribonucleotides are given in figure 2. The oligonucleotide is to be understood as a detail out of a longer nucleotide chain.

Of course, the oligonucleotides may be composed of elements of either figures.

In figures 1 and 2, lit. B means an organic base such as adenine (A), guanine (G), cytosine (C), inosine (I), uracil (U) and thymine (T) which are coupled to the deoxyribose. The linkages between the nucleotides are either phosphodiester bonds as in naturally occurring DNA or linkages spacing the nucleotides in such a way to allow hybridization with its target nucleic acid or binding to a protein in order to regulate its activity, such as e.g. phosphorothicate linkages, methylphosphonate linkages, phosphoramidate linkages or peptide linkages.

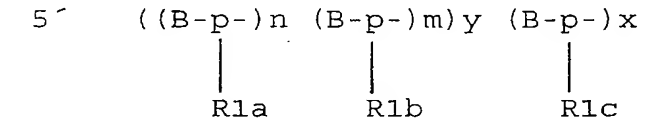
 $R_2$  and  $R_3$  represent further residues of the oligonucleotide or derivative.

 $R_4$  represents OH or a modification such as a 2'-methoxy ethoxy derivative.

The modifications of the phosphodiester linkage, shown in figures 1 and 2 can be selected from, but are not limited to.

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- 1. Oligodeoxy-ribonucleotides or oligoribionucleotides substituted by
- 1.1 R1 = 0
- 1.2 R1 = S
- 1.3. R1 = F
- 1.4. R1 =  $CH_3$
- 1.4. R1 = OEt
- 2. Oligodeoxy-ribonucleotides where R1 is varied at the internucleotide phosphates within one oligonucleotide



where lit. p stands for the phosphodiester or the phosphoramidate linkage, modified by coupling to R1a, R1b or R1c or for a peptide linkage, or for linkages spacing the nucleotides in such a way to allow hybridization with its target nucleic acid or binding to a protein in order to regulate its activity, structure, function or expression level.

where lit. B = any deoxy-ribonucleotide or ribonucleotide, depending on gene sequence according to the invention.

n, m, x, y = integers 0 - 20

Preferred maximal length of the total number of bases is 30.

2.1	$R_{1a} = S$	$R_{1b}=CH_3$	$R_{1c}=S$
2.2	$R_{1a} = S$	$R_{1b}=CH_3$	$R_{1c} = O$
2.2	$R_{1a} = S$	$R_{1b}=O$	$R_{1c}=S$
2.2	$R_{1a} = S$	$R_{1b}=O$	$R_{1c} = CH_3$
2.3	$R_{1a} = CH_3$	$R_{1b}=S$	$R_{1c} = CH_3$
2.4	$R_{1a} = CH_3$	$R_{1b}=S$	$R_{1c}=O$
2.5	$R_{1a} = CH_3$	$R_{1b}=O$	$R_{1c} = CH_3$
2.6	$R_{1a} = CH_3$	$R_{1b}=O$	$R_{1c}=S$

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2.7	$R_{1a} = O$	$R_{1b}=S$	$R_{1c}=O$
2.8	$R_{la} = 0$	$R_{1b}=S$	$R_{1c}=CH_3$
2.9	$R_{1a} = O$	$R_{1b}=CH_3$	$R_{1c}=O$
2.10	$R_{1a} = O$	$R_{1b} = CH_3$	$R_{1c}=S$

Preferably, the oligonucleotide comprises a minimum of 10 elements and a maximum of 24 elements capable of forming either 2 or 3 hydrogen bonds per element. The oligonucleotides of the invention can have modifications to the base, the sugar or the phosphate moiety. Preferred modifications are phosphorothioate (S-ODN)internucleotide linkages, and/or methylphosphonate internucleotide linkages, N'3 -> P5' phosphoramidate linkages, peptide linkages or 2'methoxyethoxy modifications of the sugar or modifications of the bases. In a very preferred embodiment the antisense oligonucleotides comprise the sequences 41 to 73, 74 to 106, 154 to 172, 173 to 203, 298 to 380, 476 to 506, 519 to 556 and 597 to 641 of figure 3 and 1273 - 1764 of figure 5. A further aspect of the invention is the use of the oligonucleotides of the invention for the inhibition of the genes p53, rb, junD, junB, TGF-S1, TGF-S2 to influence cell proliferation, in particular of primary cell cultures such as liver cells, kidney cells, osteoclasts, osteoblasts and/or keratinocytes and/or cells of the blood lineage, such as bone marrow stem cells, and/or progenitor cells of red and white blood cells and/or organ stem cells.

The Sequences 41 - 73 and/or 74 - 106 and/or 154 - 203 and/or 519 - 556 and/or 597 - 641 and/or 1273 - 1277 and/or 1481 - 1490 and/or 1532 - 1549 and/or 1656 are useful for the treatment and/or prevention of immunosuppressive disorders including, but not limited to immunosuppression in neoplastic diseases - including gliomas and other brain tumors, sarcomas, carcinomas and lymphomas - and/or immunosuppression as side effect from drugs, including, but not limited to side effects from cytotoxic agents and/or immunosuppression in AIDS patients.

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In a further embodiment of the invention these sequences are also useful for the treatment and/or prevention of hyoproliferation of normal cells, including, but not limited to immune cells, bone marrow stem cells, endothelial cells, organ stem cells and proliferating cells of the intestine.

The Sequences 41 - 73 and/or 74 - 106 and/or 298 - 380 and/or 476 - 506 and/or 519 - 556 and/or 1273 - 1480 and/or 1596 - 1614 and/or 1657 - 1658 and/or 1690 and/or 1696 - 1712 and/or 1751 and/or 1753 - 1754 and/or 1757 are useful for the treatment and/or prevention of hyperproliferative disorders, including but not limited to brain tumors, sarcomas, carcinomas and lymphomas, restenosis, hyperplasisa, pulmonary fibrosis, angiogenesis and psoriasis.

The Sequences 1278 - 1480 and/or 1491 - 1531 and/or 1582 - 1595 and/or 1615 - 1655 and/or 1691 - 1694 and/or 1697 - 1750 and/or 1759 - 1764 are useful for the treatment and/or prevention of diseases characterised by hyperfunction of the immune system and/or of inflammatory disorders and/or auto-immune disorders, including, but not limited to asthma (molecules according to the invention being applied by inhalation and/or by parenteral routes and/or orally), multiple sclerosis, inflammatory disorders of the intestine, including jejunitis, ileitis and/or colitis, as well as inflammatory disorders characterised by hyperproliferation and/or hyperfunction of cells of the eosinophilic lineage and/or glomerulonephritis and/or rejection of transplants.

The Sequences 476 - 506 and/or 1550 - 1581 and/or 1582 - 1595 and/or 1658 - 1689 and/or 1691 - 1694 and/or 1713 - 1752 are useful for the treatment and/or prevention of diseases associated with cell degeneration, including, but not limited to neurodegeneration, e.g. Alzheimer's diseases, Parkinson's, ischemic disorders, including myocardial ischemia and/or ischemia of the nervous system, including stroke.

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A further aspect of the present invention is a medicament comprising an oligonucleotide according to the invention together with additives. The oligonucleotides of the invention can be used for the preparation of a medicament for the prevention or the treatment of neoplasm, hypoproliferation, hyperproliferation, degenerative diseases, neurodegenerative diseases, ischaemia, disorders of the immune system and/or infectious diseases and can be used for the analysis of gene function or drug target validation.

Molecules according to the invention can be used to study the function of target molecules and their encoded transcription and/or translation products, including RNA molecules and proteins. Downregulations of a protein or nucleic acid molecule using molecules according to the invention can be used to study the function of the molecule. It is also a feature of the invention that molecules according to the invention can be used to study whether modulation of the product has a desired effect, including therapeutic effects and to use this information to develop a different molecule, in order to modulate the function of the protein.

This includes, for example, drug target validation with a molecule according to the invention, in order to answer the question whether development of an agent capable of modulating the structure, function or expression of a potential target molecule, e. g. an agonist or antagonist of the target molecule has desired effect and may e. g. be of therapeutic or diagnostic use.

It is thus also a feature of the invention that molecules according to the invention can be used for drug target validation, including but not limited to studying whether modulation of a protein or nucleic acid molecule has a desired effect, including therapeutic effects and using this information to develop a compound, e. g. a therapeutic compound capable of modulating the structure, function or

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expression of the molecule the function of which was previously studied with molecules according to the invention.

## Example 1

Treatment of Peripheral blood mononuclear cells with TGF-ß1 antisense phosphorothioate oligodeoxynucleotides:

Human peripheral blood mononuclear cells (PBMCs) produce transforming growth factor £1 (TGF-£1). The TGF-£1 produced by these cells negatively regulates immune cell proliferation in an autologous manner. This autologous negative regulation of immune cell proliferation could be reversed by antisense TGF-£1 molecules according to the invention, leading to stimulation of immune cell proliferation. In contrast to the molecules according to the invention, antisense molecules chosen conventionally, including that published by Hatzfeld et al. (1991) did not stimulate immune cell proliferation. Even more surprising, several sequences, chosen conventionally, even reduced immune cell proliferation.

Peripheral blood mononuclear cells (PBMCs) were isolated from venous blood of healthy donors by mixing with an equal volume of RPMI 1640 medium (Gibco) supplemented with 10 % fetal calf serum and 1 mM L-glutamine, followed by layering onto Ficoll-Hypaque (Pharmacia) gradients and centrifugation at 400 g for 30 min. PBMCs were removed from the plasma-Ficoll interface and washed in the above medium. Cells (2 x 104 in 100  $\mu$ l of medium) were plated into 96 well flat-bottom microtiter plates (Nunc) in serum supplemented complete medium. Cells were activated with 3  $\mu$ g/ml phytohemagglutinin and incubated with either no oligodeoxynucleotide (untreated control cells) or with 8 µM of different antisense phosphorothicate oligodeoxynucleotides, complementary to different regions of the human TGF-S1 mRNA for 4 days. Cells were then stained with trypan blue to determine cell viability and counted in a Neubauer counting chamber.

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Oligonucleotide sequences were either 33 sequences according to the invention, named sequences TGF-ß1-1 - TGF-ß1-33 or the TGF-ß1 antisense sequence from Hatzfeld et al. (1991), J. Exp. Med., 174, pp. 925 - 929 or 39 other conventionally chosen antisense sequences complementary to human TGF-ß1 mRNA, named N1 - N39 (see figure 3).

Surprisingly the molecules according to the invention were much more effective than antisense TGF-ß1 molecules that were chosen conventionally.

Sequences TGF-ß1-1 - TGF-ß1-33 (see figure 3) enhanced lymphocyte proliferation to between 135 and 213% of untreated controls. In contrast, treatment with the antisense sequence from document Hatzfeld et al. reduced proliferation to 62,8%.

Cells treated with the conventionally chosen TGF-ß1 antisense sequences N1 - N39 surprisingly not only failed to increase lymphocyte proliferation, but several of these sequences even revealed a marked inhibition of cell proliferation to between 51,4% and 77% of controls (sequences N1- N14, N20, N26 and N30 - N39). The antisense TGF-ß1 sequences N15 - N19, N21 - N25, N28 and N29 showed neither significant enhancement nor significant inhibition of cell proliferation with values between 94% and 103%. Sequence N27 showed slight toxicity with a reduction in cell proliferation to 88%.

Inhibition of cell proliferation by some of the TGF-ßl sequences suggests that they may not be merely ineffective, but also toxic. Analysis of the 26 sequences N1- N14, N20, N26 and N30 - N39 revealed that 23 of them contained either 2 or more sequence motifs with three consecutive Gs (hereafter called GGG motif) or at least one motif with 4, 5, or 6 Gs (motifs GGGG, GGGGG, or GGGGGG). Analysis of the sequence from Hatzfeld et al., which also inhibited PBMC proliferation, surprisingly showed that it too contains a GGGGGG plus a GGG motif. The 3 toxic sequences that contained

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neither 2 GGG motifs nor a motif of 4 or more consecutive Gs, i.e. sequences N8, N26, and N35 were found have a base content with 11 - 13 G-bases per sequence.

In contrast to the sequences from Hatzfeld et al., N1- N14, N20, N26 and N30 - N39 the sequences TGF-ß1-1 - TGF-ß1-33 showed a G-content of maximally 6 G-bases, no combination of two GGG motifs within a single sequence and no GGGG, GGGGG or GGGGGG motif. Since the TGF-ß1 mRNA contains more than 85 target regions for a GGG antisense motif and more than 34 target regions for a GGGG antisense motif, this finding in the sequences according to the invention was highly unlikely on a statistical basis.

The non-effective sequences N15 - N19, N21 - N25, N28 and N29 were found to contain a different base content from both the toxic and the effective sequences: They content of the bases A and T taken together (A/T-content) ranged from 14,3% to 28,5%. These sequences neither enhanced nor did they inhibit PBMC proliferation. Thus, they appeared to be neither effective nor toxic. In contrast to these non-effective sequences with an A/T content of 14,3% - 28,5%, the effective sequences TGF-B1-1 - TGF-B1-33 were found to have an A/T content of between 33% - 71,4%.

A further difference between the sequences of the invention and two thirds of the other sequences was found with respect to non-specific protein binding: Sequences from document Hatzfeld et al. and N1- N14, N20, N26 and N30 - N39 were found to show markedly enhanced non-specific protein binding compared to the sequences of the invention.

Sequences from Hatzfeld et al. (H) and N1 - N39 are shown in figure 3 as well as TGF-ß1 antisense sequences according to the invention.

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The finding that, while the sequences TGF-ß1-1 - TGF-ß1-33 stimulated proliferation of PBMC immune cells, the sequence from Hatzfeld et al. and sequences N1- N39 where either non-effective with little alteration in PBMC proliferation or had toxic effects and inhibited PBMC proliferation was extended to further antisense sequences both of TGF-ß2 and other genes as detailed in the following examples 2 - 7.

The sequences of the oligonucleotides related with TGF-ß1 are listed in figure 3 for the sake of ease of readability.

For certain applications, including, but not limited to application in dividing cells, including tumor cells, nucleic acid or functional or structural analogs or effective derivatives thereof according to the invention were coupled to folic acid, either at one of the carboxy-groups or at one of the nitrogen atoms of the folic acid.

Furthermore, for certain applications, nucleic acid or functional or structural analogs or effective derivatives thereof according to the invention are mixed with and/or coupled to hormones, steroid hormones such as oestrogen, progesterone, corticosteroids, mineralocorticoids, androgens, phospholipids, peptides, proteoglycans, glycolipids and derivatives therefrom. Preferably, a coupling occurs at R<sup>2</sup> and/or R<sup>3</sup> of figures 1 and 2.

#### Example 2

p53 antisense nucleic acids (figure 3 shows the respective oligonucleotides)

p53 is a tumor suppressor gene that negatively regulates cell proliferation. Certain mutations in the gene can alter the function of p53 in such a way that it becomes an oncogene. The effects of p53 antisense oligodeoxynucleotides on cells

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containing wild type p53 was analyzed and subsequently also the effect of these sequences on cells with mutated p53.

In cells with wild type p53 effective antisense nucleic acids will lead to downregulation of the wild type p53 protein and thus to enhanced proliferation of the treated cells. Molecules according to the invention are named p53-1 - p53-33. Noneffective p53 antisense sequences were named p53-N-1 - p53-N-18. Toxic sequences, which inhibited proliferation instead of enhancing it as do effective p53 antisense sequences were named p53-T-1 - p53-T-29.

Normal human fibroblasts were grown in RPMI medium supplemented with 5% fetal calf serum (FCS) and 2500 cell/well were plated into 96-well microtiter plates. Antisense phosphorothioate oligonucleotides were added at 2  $\mu$ M concentration after 2 h.

Two assays to determine cell proliferation were performed:

- To determine 3H-thymidine incorporation, cells were incubated before harvesting with 0,15  $\mu$ Ci 3H-thymidine/well for 6 h. Cells were lysed by freezing, spotted onto glass filters and the amount of incorporated tritium was determined by liquid scintillation counting.
- To determine cell number, cells were stained with trypan blue and counted in a Neubauer counting chamber.

Surprisingly, only treatment of cells with antisense sequences according to the invention (p53-1 - p53-33) resulted in an increase in thymidine incorporation to between 3- and 9-fold.

In contrast, treatment with noneffective sequences (p53-N-1 - p53-N-18) did not result in significant alterations in thymidine incorporation.

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Furthermore, treatment with toxic antisense p53 sequences (p53-T-1- p53-T-29) resulted in a decrease in proliferation instead of an increase.

In summary, the 33 antisense sequences according to the invention resulted in effective downregulation of negative growth control by p53 and increased cell proliferation, while the 47 other antisense sequences had either no significant effect on cell proliferation or even suppressed cell proliferation.

Example 3

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junB antisense nucleic acids (figure 3 shows the respective oligonucleotides)

junB and junD, two genes encoding transcription factors of the jun gene family are negative regulators of cell growth, like p53. The effects of different junB and junD antisense oligodeoxynucleotides was analyzed.

Effective junB and JunD antisense nucleic acids will lead to downregulation of the JunB an JunD proteins respectively and thus to enhanced proliferation of the treated cells. Antisense molecules according to the invention are named JunB-1 - JunB-19 and JunD-1 - JunD-31. Noneffective junB antisense sequences were named JunB-N-1 - JunB-N-57. Toxic sequences, which inhibited proliferation instead of enhancing it were named JunB-T-1- JunB-T-20 and JunD-T-1 - JunD-T-17.

Normal human fibroblasts were grown in RPMI medium supplemented with 5% fetal calf serum (FCS) and 2500 cell/well were plated into 96-well microtiter plates. Antisense phosphorothioate oligonucleotides were added at 2  $\mu$ M concentration after 2 h.

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Two assays to determine cell proliferation were performed:

- To determine 3H-thymidine incorporation, cells were incubated before harvesting with 0,15  $\mu$ Ci 3H-thymidine/well for 6 h. Cells were lysed by freezing, spotted onto glass filters and the amount of incorporated tritium was determined by liquid scintillation counting.
- To determine cell number, cells were stained with trypan blue and counted in a Neubauer counting chamber.

Surprisingly, again only treatment of cells with antisense sequences according to the invention (JunB-1 - JunB-19 and JunD1- JunD31) resulted in an increase in thymidine incorporation to between 2- and 7-fold.

In contrast, treatment with noneffective sequences (JunB-N-1 - JunB-N-57) did not result in significant alterations in thymidine incorporation.

Furthermore, treatment with toxic antisense junB or JunD sequences (JunB-T-1- JunB-T-20 and JunD-T-1 - JunD-T-17) resulted in a decrease in proliferation instead of an increase.

In summary, the 50 antisense sequences according to the invention resulted in effective downregulation of negative growth control by JunB and JunD, while the 94 other antisense sequences had either no significant effect on cell proliferation or were even toxic.

Example 4 (figure 3 shows the respective oligonucleotides)

erbB-2, is a transmembrane molecule with an intracellular tyrosine kinase activity that is amplified and/or overex-pressed by carcinoma cells in a variety of neoplasms including breast cancer, lung cancer, oesophageal and gastric

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cancer, bile duct carcinoma, bladder cancer, pancreatic cancer and ovarian cancer.

In several of these tumors, an amplification and overexpression of the c-erbB-2 gene in the tumor tissue has been shown to correlate with a poor clinical prognosis. Overexpression of p185erbB-2 in non-small-cell lung carcinoma has been shown to impart resistance to a number of chemotherapeutic agents.

Effective erbB-2 antisense nucleic acids will lead to downregulation of the erbB-2 protein and in overexpressing tumor cell lines will lead to reduced cell proliferation of the treated cells. Antisense molecules according to the invention are named erbB-2-1 - erbB-2-83. Noneffective erbB-2 antisense sequences were named erbB-2-N-1 - erbB-2-N-95.

erbB-2 overexpressing SK-Br-3 human mammary carcinoma cells were grown in RPMI medium supplemented with 5% fetal calf serum (FCS) and 2500 cell/well were plated into 96-well microtiter plates. Antisense phosphorothioate oligonucleotides were added at 2  $\mu$ M concentration after 2 h.

To determine erbB-2 protein expression cells were harvested with a cell scraper and subjected to ELISA protein determination.

Only treatment of cells with antisense sequences according to the invention (erbB-2-1 - erbB-2-83) resulted in a significant reduction in erbB-2 protein expression by 40-95%.

In contrast, treatment with noneffective sequences (erbB-2-N-1 - erbB-2-N-95) did not result in significant alterations in erbB-2 protein expression.

To determine cell number, cells were stained with trypan blue and counted in a Neubauer counting chamber.

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Only treatment of cells with antisense sequences according to the invention (erbB-2-1 - erbB-2-83) resulted in a reduction in cell number by 35-70%.

In contrast, treatment with noneffective sequences (erbB-2-N-1 - erbB-2-N-95) did not result in significant alterations in cell proliferation.

erbB-2 antisense sequences were shown in figure 3-8 to 3-11

Example 5 (figure 3 shows the respective oligonucleotides)

The c-fos gene encodes an immediate early gene type transcription factor. Effective c-fos antisense nucleic acids will lead to downregulation of the c-Fos protein.

Antisense molecules according to the invention are named c-fos-1 - c-fos-31. Noneffective c-fos antisense sequences were named c-fos-N-1 - c-fos-N-12.

Normal human fibroblasts were grown in RPMI medium supplemented with 5% fetal calf serum (FCS) and 2500 cell/well were plated into 96-well microtiter plates. Antisense phosphorothioate oligonucleotides were added at 2  $\mu$ M concentration after 2 h.

Expression of the c-Fos protein was determined by ELISA in cell lysates.

Only treatment of cells with antisense sequences according to the invention (c-fos-1 - c-fos-31) resulted in a significant reduction in c-fos protein expression by 45-95%.

In contrast, treatment with noneffective sequences (c-fos-N-1 - c-fos-N-12) did not result in significant alterations in c-Fos protein expression.

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Example 6 (figure 3 shows the respective oligonucleotides)

TGF-ß2, like TGF-ß1 is a member of the transforming growth factor-ß family of cytokines.

Overexpression of TGF-ß1 and TGF-ß2 is linked to malignant progression, immunosuppression and escape of the tumors from surveillance by the immune system.

Effective TGF-S2 antisense nucleic acids will lead to downregulation of the TGF-S2 growth factor.

Antisense molecules according to the invention are named TGFß2-1 - TGF-ß2-38. Noneffective TGF-ß2 antisense sequences were named TGF-ß2-N-1 - TGF-ß2-N-40.

TGF-ß2 overexpressing tumor cells were grown in RPMI medium supplemented with 5% fetal calf serum (FCS) and 2500 cell/well were plated into 96-well microtiter plates. Antisense phosphorothioate oligonucleotides were added at 2  $\mu$ M concentration after 2 h.

TGF-ß2 protein expression was determined by ELISA, both in the supernatant and in cell lysates.

Only treatment of cells with antisense sequences according to the invention (TGF-ß2-1 - TGF-ß2-38) resulted in a significant reduction in TGF-ß2 protein expression by 35-80%.

In contrast, treatment with noneffective sequences (TGF-ß2-N-1 - TGF-ß2-N-40) did not result in significant alterations in TGF-ß2 protein expression.

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Example 7 (figure 3 shows the respective oligonucleotides)

rb antisense nucleic acids

rb is a tumor suppressor gene that negatively regulates cell proliferation. The effects of rb antisense oligodeoxynucleotides on cells containing wild type rb was analyzed.

In cells with wild type rb effective antisense nucleic acids will lead to downregulation of the wild type rb protein and thus to enhanced proliferation of the treated cells. Molecules according to the invention are named rb-1 - rb-45. Noneffective rb antisense sequences were named -1 - rb-N-168. Toxic sequences, which inhibited proliferation instead of enhancing it as do effective rb antisense sequences were named rb-T-1- rb-T-16.

Normal human fibroblasts were grown in RPMI medium supplemented with 5% fetal calf serum (FCS) and 2500 cell/well were plated into 96-well microtiter plates. Antisense phosphorothicate oligonucleotides were added at 2  $\mu$ M concentration after 2 h.

Two assays to determine cell proliferation were performed:

- To determine 3H-thymidine incorporation, cells were incubated before harvesting with 0,15  $\mu$ Ci 3H-thymidine/well for 6 h. Cells were lysed by freezing, spotted onto glass filters and the amount of incorporated tritium was determined by liquid scintillation counting.
- To determine cell number, cells were stained with trypan blue and counted in a Neubauer counting chamber.

Surprisingly, only treatment of cells with antisense sequences according to the invention (rb-1 - rb-45) resulted in an increase in thymidine incorporation to between 2- and 6-fold.

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In contrast, treatment with noneffective sequences (rb-N-1 - rb-N-168) did not result in significant alterations in thymidine incorporation.

Furthermore, treatment with toxic antisense rb sequences (rb-T-1-rb-T-16) resulted in a decrease in proliferation instead of an increase.

In summary, the 45 antisense sequences according to the invention resulted in effective downregulation of negative growth control by rb and increased cell proliferation, while the 184 other antisense sequences had either no significant effect on cell proliferation or even suppressed cell proliferation.

#### Example 8

Oligonucleotide sequences according to the invention were synthesized with various different backbone modifications: Exemplary results are given below.

For the sequence

erbB-2-42: CATCTGGAAACTTCCAGATG

the following chemical modifications were tested in erbB-2 overexpressing carcinoma cells:

1. S-ODN erbB-2-42 (i.e. all backbone linkages were thioate modifications).

2. Me-ODN/S-ODN/Me-ODN erbB-2-42 (i.e. Linkages at the 5'and 3'end were methylphosphonate linkages while linkages in the middle were thioate modifications as follows):

C-pMe-A-pMe-T-pS-C-pS-T-pS-G-pS-G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pS-C-pS-A-pS-G-pS-A-pMe-G or

C-pMe-A-pMe-T-pMe-C-pS-T-pS-G-pS-G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pS-C-pS-A-pS-G-pMe-A-pMe-T-pMe-G or

C-pMe-A-pMe-T-pMe-C-pMe-T-pS-G-pS-G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pS-C-pS-A-pMe -G-pMe-A-pMe-T-pMe-G or

C-pMe-A-pMe-T-pMe-C-pMe-T-pMe-G-pMe-G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pMe-C-pMe-A-pMe-G-pMe-A-pMe-T-pMe-G

3. Me-ODN / S-ODN erbB-2-42 (i.e. Linkages at the 5' end were methylphosphonate linkages while linkages at the 3' were thioate modifications as follows):

C-pMe-A-pMe-T-pMe-C-pMe-T-pMe-G-pMe-G-pMe-A-pMe-A-pMe-A-pS-C-pS-T-pS-T-pS-C-pS-C-pS-A-pS-G-pS-A-pS-T-pS-G

4. S-ODN / Me-ODN erbB-2-42 (i.e. Linkages at the 5 end were methylphosphonate linkages while linkages at the 3 were thioate modifications as follows):

C-pS-A-pS-T-pS-C-pS-T-pS-G-pS-G-pS-A-pS-A-pS-A-pMe-C-pMe-T-pMe-T-pMe-C-pMe-C-pMe-A-pMe-G-pMe-A-pMe-T-pMe-G

5. Me-ODN erbB-2-42 (i.e. linkages methylphosphonate linkages):

C-pMe-A-pMe-T-pMe-C-pMe-T-pMe-G-pMe-G-pMe-A-pMe-A-pMe-A-C-pMe-T-pMe-T-pMe-C-pMe-C-pMe-A-pMe-G-pMe-A-pMe-T-pMe-G-pMe-G-pMe-A-pMe-G-pM

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6. pN/S-ODN/pN erbB-2-42 (i.e. Linkages at the 5'and 3'end were phosphoramidate linkages while linkages in the middle were thioate modifications as follows):

C-pN-A-pN-T-pS -C-pS-T-pS-G-pS-G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pS-C-pS-A-pS-G-pS-A-pN-T-pN-G

or

C-pN-A-pN-T-pN-C-pS-T-pS-G-pS-G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pS-C-pS-A-pS-G-pN-A-pN-T-pN-G

or

C-pN-A-pN-T-pN-C-pN -T-pS-G-pS-G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pS-C-pS-A-pN -G-pN-A-pN-T-pN-G

or

C-pN-A-pN-T-pN-C-pN -T-pN -G-pN -G-pS-A-pS-A-pS-A-pS-C-pS-T-pS-T-pS-C-pN -C-pN-A-pN -G-pN-A-pN-T-pN-G

#### where

pS stands for substitution of one of the non-bridging oxygen atoms of the backbone linkage with a sulfur atom, while pMe stands for substitution of one of the non-bridging oxygen atoms of the backbone linkage with a methyl group. pN stands for a N3´->P5´ phosphoramidate linkage.

Also a combination of linkages  $(N-pS-N-pO-N-pO-N)_n-[pS-N]_m$  wherein n=1-10 and m=0-6 where N stand for any nucleotide or structural or functional analog or derivative thereof.

While the Me-ODN backbone modification strongly reduced the erbB-2 activity of the erbB-2-42 sequence to less than 20%, backbone modifications 1.-4. had strong erbB-2 inhibitory capacity with an inhibition of erbB-2 protein expression by between 78% and 89% at 2  $\mu$ M concentration at 48 h after the beginning of treatment of overexpressing carcinoma cells. While the pure S-ODN had the highest suppression capacity with 89%, the Me-ODN/S-ODN/Me-ODN as well as the Me-ODN/S-ODN

and S-ODN/Me-ODN and pN/S-ODN/pN, displayed reduced protein binding and when tested for complement activation, showed reduced complement activation. These characteristics are advantageous for certain applications e.g. intravenous systemic application in vivo.

# Example 9

Similar effects were obtained when testing other sequences according to the invention with the above backbone modifications.

Inhibition of TGF-beta-1 gene expression with the effective sequences for TGF-beta-1 according to the invention was highest with S-ODN and the Me-ODN/S-ODN/Me-ODN backbone modifications and lowest with the Me-ODN modification, while protein binding and complement activation were reduced in sequences containing Me-ODN linkages.

#### Example 10

Surprisingly, effectivity of sequences according to the invention was significantly improved in various cell types by coupling nucleic acids according to the invention to folic acid:

erbB-2 inhibitory capacity which was relatively low after 24 h compared to 48 h with an inhibition of erbB-2 protein synthesis by 24-37% was markedly increased by coupling sequences according to the invention to folic acid to 48-62% at 2  $\mu$ M concentration 24 h after the beginning of treatment of overexpressing carcinoma cells.

Similar effects were achieved by coupling sequences according to the invention to folic acid derivatives including aminopterin and amethopterin.

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### Example 11

Surprisingly, effectivity of sequences according to the invention was strongly improved by coupling oligonucleotides according to the invention to cortisol:

Cellular uptake and inhibitory capacity of sequences according to the invention including sequences for TGF-beta-1, TGF-beta-2, c-fos, p53, erbB-2, rb, c-fos, junB, junD, c-jun, MIP-1 alpha, JAK-2, bcl-2 and were markedly increased by coupling cortisol either to the 3 or 5 hydroxyl groups of oligonucleotide sequences according to the invention.

### Example 12

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Effectivity of sequences according to the invention was also strongly improved in various cell types by coupling nucleic acids according to the invention to or mixing them with other steroid hormones and their derivatives, including oestrogens, anti-oestrogens, prednisone, prednisolone, androgens, anti-androgens, gestagenes like progesterone as well as peptides, proteoglycans, glycolipids, phospholipids and derivatives therefrom.

Androgens, particularly androstendion and testosterone, as well as anti-androgens, including cyproteronacetate, flutamide, anandrone, linked to the nucleic acids increased effectiveness of the molecules in various cell types including prostatic carcinoma cells.

Oestrogens, anti-oestrogens and their derivatives, including fosfestrol, toremifen, ethinyloestradiole, diethylstilboestole and the oestradiole derivatives oestradiol-benzoate, oestradiol-valerinate and oestradiol-undecylate, as well as progesterone and its derivatives, including medroxyprogestroneacetate and megestrolacetate linked to the oligonucleotides strongly enhanced activity of the molecules according

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to the invention in various cell types including mammary carcinoma cells.

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#### Claims

 A method for the preparation of an antisense oligonucleotide or derivative thereof comprising the steps of

- selecting a target nucleic acid, if necessary elucidating its sequence
- generating the antisense oligonucleotide with the proviso that
  - the oligonucleotide comprises at least 8 residues,
  - the oligonucleotide comprises at maximum twelve elements, which are capable of forming three hydrogen bonds each to cytosine bases,
  - the oligonucleotide does not contain four or more consecutive elements, capable of forming three hydrogen bonds each with four consecutive cytosine bases (CCCC) within the target molecule or alternatively four or more consecutive elements of GGGG,
  - the oligonucleotide does also not contain 2 or more series of three consecutive elements, capable of forming three hydrogen bonds each with three consecutive cytosine bases (CCC) within the target molecule, or alternatively 2 or more series of three consecutive elements of GGG, and
  - the ratio between residues forming two hydrogen bonds per residue (2H-bond-R) with the target molecule and those residues forming three hydrogen bonds per residue (3H-bond-R) with the target molecule, is ruled by the following specifications:

3H-bond-R

 $\geq 0.29$ 

3H-bond-R + 2H-bond-R

and synthesizing the oligonucleotide thus generated in a per se known manner.

2. The method according to claim 1, wherein the generated oligonucleotide complies with the following specification

3H-bond-R = 0.33 to 0.86 3H-bond-R + 2H-bond-R

- 3. The method according to any one of the claims 1 or 2, wherein the generated oligonucleotides are modified for higher nuclease resistance than naturally occurring oligo- or polynucleotides.
- 4. The method according to claim 3, wherein the generated oligonucleotides are modified at the bases, the sugars or the linkages of the oligonucleotides, preferably by phosphorothicate (S-ODN) internucleotide linkages, and/or methylphosphonate internucleotide linkages, N'3 -> P5' phosphoramidate linkages, peptide linkages or 2'-methoxyethoxy modifications of the sugar or modifications of the bases.
- 5. The method according to claim 3 and/or 4, wherein the oligonucleotide has at least two different types of modifications.
- 6. The method according to any one of the claims 1 to 5, wherein the oligonucleotides are reacted with folic acid, hormones such as steroid hormones or corticosteroides or derivatives thereof by linking the oligonucleotides covalently to or mixing with folic acid, hormones such as steroide hormones or corticosteroides, peptides, proteoglycans, glycolipids or phospholipids.

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- 7. An antisense oligonucleotide or derivative thereof obtainable according to the method according to any one of the claims 1 to 6 except oligonucleotides represented by Fig. 4.
- 8. The oligonucleotide or derivative of claim 7, which does not contain four or more consecutive guanosine  $(N_a GGGGN_b)$  or inosine  $(N_a IIIIN_b)$  residues and the oligonucleotide does not contain two or more series of three or more consecutive guanosine residues  $(N_a GGGN_c GGGN_b)$  and does not contain two ore more series of three or more consecutive inosine residues  $(N_a IIIN_c IIIN_b)$ , wherein  $N_a$ ,  $N_b$ ,  $N_c$  represent indepently nucloetides or oligonucleotides or derivatives thereof having 0 to 20 residues.
- 9. The oligonucleotide or derivative of claims 7 and/or 8, comprising a minimum of ten elements and a maximum of 24 elements capable of forming either two or three hydrogen bonds per element.
- 10. The oligonucleotide or derivative according to any one of the claims 7 to 9, having modifications at the bases, the sugars or the phosphate moieties of the oligonucleotides.
- 11. The oligonucleotide or derivative of any one of the claims 7 to 10, wherein the modifications are phosphorothicate (S-ODN) internucleotide linkages, and/or methylphosphonate internucleotide linkages, N'3 -> P5' phosphoramidate linkages, peptide linkages or 2'-methoxyethoxy modifications of the sugar or modifications of the bases.

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- 12. The oligonucleotide or derivative of any one of the claims 7 to 11 coupled to or mixed with folic acid, hormones, steroid hormones such as oestrogene, progesterone, corticosteroids, mineral corticoids, peptides, proteoglycans, glycolipids, phospholipids and derivatives therefrom.
- The oligonucleotide according to any one of the claims 13. 7 to 12, wherein the antisense oligonucleotide against the TGF-S1 gene comprise the sequences 41 to 73 of Fig. 3, the oligonucleotides against the gene p53 comprising the sequences 74 to 106 of Fig. 3, the antisense oligonucleotides against junB comprising the sequences 154 to 172 of Fig. 3, the antisense oligonucleotides against junD comprising the sequences 173 to 203 of Fig. 3, the antisense oligonucleotides against the erbB-2 gene comprise the sequences 298 to 380 of Fig 3, the antisense oligonucleotides against c-fos genes comprise the sequences 476 - 506 of Fig. 3; the antisense oligonucleotides against the gene TGF-S2 comprise the sequences 519 to 556 of Fig. 3 as well as the antisense oligonucleotides against the gene rb comprise the sequences 597 to 641 of Fig. 3.; as well as sequences 1273 to 1764. of Fig. 5.
- 14. A composition comprising an oligonucleotide or derivative according to any one of the claims 7 to 13 for the manufacturing of a medicament or a composition for the inhibition of the genes p53, rb, junD, junB, TGF-B1, TGF-B2 to influence cell proliferation, in particular of primary cell cultures such as liver cells, kidney cells, osteoclasts, osteoblasts and/or keratinocytes and/or cells of the blood lineage, such as bone marrow stem cells, and/or progenitor cells of red and white blood cells.

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- 15. A medicament comprising an oligonucleotide according to any one of the claims 7 to 13 together with additives.
- 16. The use of the oligonucleotides according to any of the claims 7 to 13 for the preparation of a medicament for the prevention or the treatment of neoplasm, hypoproliferation, hyperproliferation, degenerative diseases, neurodegenerative diseases, ischaemia, disorders of the immune system and/or infectious diseases, and/or metabolic dysfunctions.
- 17. The use of the oligonucleotides according to any one of the claims 7 to 13 for the analysis of gene function or drug target validation.

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FIG. 2

		•	
1.		A3	CCCGGAGGCGGCATGGGGGA
2.		Nl	CCTCAGGGAGAAGGGCGC
3.		N2	GTAGGAGGCCTCGAGGG
4.		N3	CTGCAGGGGTTGGGGGTC
5.		N4	AGGGCTGGTGTGGGG
		_ · · ·	
6.		N5	GGCATGGGGGAGGCGCG
7.		N6	CCGGAGGCGCATGGGG
8.		N7	GGGGGCTGGCGAGCCGC
9.		N8	GGACAGGATCTGGCCGCGGATGG
10.		N9	CCCCTGGCTCGGGGGC
11.		N10	GGGCCGGCGCACCTCC
12.		N11	GGGCAGCGGCCGGGCGG
			ACGGCCTCGGGCAGCGGG
13.		N12	
14.		N13	GGGTGCTGTTGTACAGGG
15.		N14	GGGTTTCCACCATTAGCACGCGGG
16.		N15	TCATAGATTTCGTT
17.		N16	TTGTCATAGATTT
18.		N17	AAGAACATATATATG
19.		N18	AAGAACATATATAT
20.		N19	TTGAAGAACATATATA
21.		N20	CCGGGAGAGCAACACGGG
22.		N21	ACTTTAACTTGA
23.		N22	ATTGTTGCTGTATTT
24.		N23	ATTGTTGCTGTATT
25.		N24	AATTGTTGCTGTATT
26.		N25	AATTGTTGCTGTAT
27.		N26	GGCGAGTCGCTGGGTGCCAGCAGCCGG
28.		N27	GGCGAGTCGCTGGG
			ACATCAAAAGATAA
29.		N28	
30.		N29	TGACATCAAAAGAT
31.		И30	GGGCCCTCTCCAGCGGGG
32.		N3I	GGGCTCGGCGGTGCCGGG
33.		N32	GGGGCAGGCCCGAGGCA
34.		N33	GGCTCCAAATGTAGGGGC
35.		N34	CGGGTTATGCTGGTTGTACAGGGC
36.		N35	CGGCGCCGAGGCGCCCGGG
37.		N36	GGGCGGGCGGACC
			GGGCGGGCGGGGGG
38.		N37	* · ·
39.		N38	GGGCGGGTGGGGCCGGG
40.		N39	GGGCAAGGCAGCGGGGGGG
41.		TGF-ß1-1	CGGTAGCAGCG
42.		TGF-£1-2	CCAGTAGCCACAGC
		TGF-B1-3	GCAGGTGGATAGTCC
43.			
44.		TGF-131-4	CTTGCAGGTGGATAG
45.		<b>TGF-ß1</b> -5	CGATAGTCTTGCAGG
46.		TGF- <b>ß1</b> -6	CCATGTCGATAGTCTTGC
47.		TGF-\$1-7	CTCGATGCGCTTCCG
48.		TGF-ß1-8	CCTCGATGCGCTTCC
49.		TGF-61-9	GGATGGCCTCGATGC
50.		TGF-B1-10	GGACAGGATCTGGCC
			CGCAGCTTGGACAGG
51.		TGF-ß1-11	
52.		TGF-£1-12	GAGCCGCAGCTTGG
53.		TGF-£1-13	CGAGCCGCAGCTTG
54.		TGF-ß1-14	ACCTCCCCTGGCT
55.		TGF-\$1-15	CCACCATTAGCACG
56.		TGF-B1-16	GAACTTGTCATAGATTTC
<b>57</b> .		TGF-£1-17	GCTGTGTGTACTCTGC
58.		TGF-ß1-18	GCTCCACGTGCTGC
59.		TGF-£1-19	GAATTGTTGCTGTATTTC
		TGF-B1-20	GCCAGGAATTGTTGC
60.			GCCAGGAATIGITGC GTGACATCAAAAGATAAC
61.		TGF-\$1-21	
62.		TGF-\$1-22	GGCTCAACCACTGCC
63.		TGF-\$1-23	GCTGTCACAGGAGC
64.		TGF-£1-24	CCTGCTGTCACAGG
65.		TGF-£1-25	GCAGTGTGTTATCCCTGC
66.		TGF-ß1-26	GCAGTGTTTATCCC
Fi~	3 - 1		
Fig.	3 - 1		

67. 68. 69. 70. 71. 72. 73.		TGF-&1-27 TGF-&1-28 TGF-&1-29 TGF-&1-30 TGF-&1-31 TGF-&1-32	CCAGGTCACCTCGG GCCATGAATGGTGGC GCCATGAATGGTGG CCATGAGAAGCAGG GGAAGTCAATGTACAGC CCACGTAGTACACGATGG GCACTTGCAGGAGC
74. 75. 77. 78. 81. 82. 83. 84. 85. 88. 89. 91. 99. 99. 99. 99. 101. 103. 106.		p53-1 p53-2 p53-3 p53-4 p53-5 p53-6 p53-6 p53-7 p53-8 p53-10 p53-11 p53-12 p53-12 p53-14 p53-15 p53-16 p53-17 p53-18 p53-19 p53-20 p53-21 p53-22 p53-23 p53-24 p53-25 p53-25 p53-26 p53-27 p53-26 p53-27 p53-29 p53-30 p53-31 p53-32 p53-32 p53-32 p53-32 p53-32 p53-32	CCATGGCAGTGACC GGCTCCTCCATGGC GCTAGGATCTGACTGC CCTGACTCAGAGGG GGTCTGAAAATGTTTCC CCATTGCTTGGGACGG GCATCAAATCATCC CCATTGTTCAATATCG GGAGCTTCAGTGAACC GGAGCTTCATCTGGACC CCTCTGGCATTCTGG AGGGACAGAAGATG GTTTTCTGGGAAG GGTTTTCTGGGAAG GGTAGGTTTCTGG GGTAGGTTTCTGG CCAGAATGCAAGAGCC GCTAGCATTCTGG CCAGAATGCAAGAAGCC GCTGTCCCAGAATGC GCAAGTCACAGACTTGGC CCACAGCTGCACAGG GGTGTGATCACC GTCATGTGCTGTGA CGCTATCTGAGCACC CCACGGATCTCAACC GCCAGAATACC CCACGGATCTCACACC CCACGGATCTCACACC CCACGGATCTCACACC CCACGGATCTCACACC CCACGGATCTCACCC CCTCATTCACCTCTCCGC CCTTGAGTTCCAACCC CCTTTTTTTCACCTTCACCC CCTTTTTTTCACCTTCACCC CCTTTTTTTCACCTTCACCC CCTTTTTTTT
107. 108. 109. 110. 111. 112. 113. 114. 115. 116. 117. 118. 119. 120. 121. 122. 123. 124.		p53-N-1 p53-N-2 p53-N-3 p53-N-4 p53-N-5 p53-N-6 p53-N-7 p53-N-8 p53-N-9 p53-N-10 p53-N-11 p53-N-12 p53-N-12 p53-N-15 p53-N-15 p53-N-15 p53-N-16 p53-N-17 p53-N-18	AAAATGTTTCCT TGAAAATGTTTC CTGAAAATGTTT TCTGAAAATGTTT TCTGAAAATGTT TCTGAAAATGTT AAATCATCCATT TTGTTCAATATC ATTGTTCAATATC ATTGTTCAATAT CATTGTTCAATAT CATTGTTCAATAT AAAAGTGTTTCT ACATGAGTTTTTTAT AACATGAGTTTTTTAT AACATGAGTTTTTTA AACATGAGTTTTTTTA AACATGAGTTTTTTTTTA AACATGAGTTTTTTTTTT
125. 126. 127. 128. 129. Fig.	3 - 2	p53-T-1 p53-T-2 p53-T-3 p53-T-4 p53-T-5	CAGAGGGGGCTCGACGC CTGACTCAGAGGGGGCTC AGGGGGACAGAACG TTGGGACGGCAAGGGGGACAGAA TGGGACGGCAAGGGGGA
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130.	p53-T-6	GCCACGGGGGGAGCA
131.	p53-T-7	GCAGGGCCACGGGGGAG
132.	2-T-8	AGGGCCACGGGG
133.	p53-T-9	CAGGGGCCACGGG
134.	p53-T-10	GGTGCAGGGCCACG
135.	p53-T-11	TGGTGCAGGGCCGCCGG
136.	p53-T-12	GGGCTGGTGCAGGGGCC
137.	p53-T-13	AGGGGCTGGTGCAGGGG
138.	p53-T-14	GGGCTGGTGCAGGG
139.	p53-1-14 p53-T <b>-1</b> 5	GAGGGGGCTGGTGCAG
140.	p53-T-16	AGGAGGGGCTGGTG
141.	<b>-</b>	GGGCCAGGAGGGGGCTGG
142.	p53-T-17 p53-T-18	· AGGGCCAGGAGGGGCT
143.	p53-1-18 p53-T-19	GGGGCCAGGAGGGG
144.	p53-1-19 p53-T-20	CAGGGCCAGGAGGG
145.	<u></u>	TCTGGGAAGGGACAGA
145.	p53-T-21	TGAGGGCAGGGGAGTA
	p53-T-22	
147.	p53-T-23	TTGAGGGCAGGGAG
148.	p53-T-24	CGGGTGCCGGGCGGGGGG
149.	p53-T-25	CGGACGCGGGTGCCGGGGGGT
150.	p53-T-26	CGGGTGCCGGGCGG
151.	p53-T-27	GGACGCGGGTGCCGGGCG
152.	p53-T-28	TGGGGGCAGCCTCACA
153.	p53-T-29	GGTGGGGCAGCCCT
154.	JunB-1	CCATTTTAGTGCACATCCGG
155.	JunB-2	CCATTTTAGTGCACATCC
156.	JunB-3	GCTGTTCCATTTTAGTGC
157.	JunB-4	GTAGTCGTGTAGAG
158.	JunB-5	GTTTGTAGTCGTGTAG
159.	JunB-6	GTTTCAGGAGTTTGTAG
160.	JunB-7	CCAGCTCCGAAGAGG
161.	JunB-8	CGTCGTCGTGATCACG
162.	JunB-9	GGTAAAAGTACTGTCC
163.	JunB-10	GGCTTTGACAAAGCC
164.	JunB-11	CTTGTGCAGATCGTCCAG
165.	JunB-12	CGTGGTTCATCTTGTGC
166.	JunB-13	CACGTGGTTCATCTTGTG
167.	JunB-14	CCTCCTTGAAGGTGG
168.	JunB-15	CGCTCCACTTTGATGCG
169.	JunB-16	CCTTGTCCTCCAGG
170.	JunB-17	GGTACTCGACAGCC
171.	JunB-18	CTGACGTGGGTCATG
172.	JunB-19	CCGTTGCTGACGTGG
173.	JunD-1	CATCCTCCGCCTCC
173. 174.	JunD-1 JunD-2	GTTTCCATCCTCCG
175.	JunD-3	GGTGTTTCCATCCTCC
176.	JunD-4	GGTGTTTCCATCCTC
177.	JunD-5	GCTCAGCGCCTCATC
178.	JunD-6	CCTTCTTCATCATGCTGC
179.		CCTTCTTCATCATGCTG
180.	JunD-7	CCTTCTTCATCATGC
	JunD-8	<del></del>
181.	JunD-9	GCGTCCTTCTTCATCATGC
182.	JunD-10	CCTGCTCACTCAGG
183.	JunD-11	CGCAGGCTTGAGCG
184.	JunD-12	GCCAGCTTCAGCAGC
185.	JunD-13	GGTGGTGACCAGCC
186.	JunD-14	CCTCGGCGAACTCC
187.	JunD-15	GCTTGTGTAAATCC
188.	JunD-16	GGTTCTGCTTGTGTAAATCC
189.	JunD-17	GCTGCTCAGGTTCGC
190.	JunD-18	GAAGGCGACCGTCG
191.	JunD-19	CGAAGGCGACCGTC
192.	JunD-20	GCACCGTCTGTGGC
193.	JunD-21	CGTGTCCATGTCGATGG
194.	JunD-22	CGTGTCCATGTCGATG
$\mathtt{Fig}_{.}$	3 - 3	
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195.	JunD-23	GCGTGTCCATGTCG
	<del>_</del>	
196.	JunD-24	CCAGCTTGCGCTTGC
197.	JunD-25	CGCTCCAGCTTGCG
198.	JunD-26	CGTGTTCTGACTCTTGAG
199.	J <b>un</b> D-2 <b>7</b>	CGTGTTCTGACTCTTG
200.	JunD-28	GCTGTTGACGTGGC
201.	JunD-29	CGACTCAGTACGCC
202.	JunD-30	GCCATGCCCGACTC
203.	JunD-31	CCCTTGGAGGTGGC
204	T 10 NY	
204.	JunB-N-1	TTTTAGTGCACAT
205.	JunB-N-2	TGTTCCATTTTAGT
	<del>-</del>	
	JunB-N-3	AAAAAAGTGGAAG
207.	JunB-N-4	TACAAAAAAAGTG
208.	JunB-N-5	ATACAAAAAAAGT
209.	JunB-N-6	CATACAAAAAAAGT
210.	JunB-N-7	CATACAAAAAAAG
211.		
	JunB-N-8	GAAAAAAACATAC
212.	JunB-N-9	CAGAAAAAAACATAC
213.	JunB-N-10	CAGAAAAAAACAT
214.	JunB-N-11	TTCAATATGAATCG
215.	JunB-N-12	TATTCAATATGAATCG
216.	JunB-N-13	
		TATTCAATATGAATC
217.	JunB-N-14	TATTCAATATGAAT
218.	JunB-N-15	TATATTCAATATGAA
219.	JunB-N-16	TTATATTCAATATGA
220.	JunB-N-17	TATTATATTCAATATGA
221.	JunB-N-18	TTATATTCAATATG
222.	JunB-N-19	TATTATATTCAATATG
223.	JunB-N-20	ATTATATTCAATAT
224.	JunB-N-21	TATTATATTCAATAT
225.	JunB-N-22	ATATATTATATTCAATAT
226.	JunB-N-23	AAATATATTATATTCAATAT
227.	JunB-N-24	TATTATATTCAATA
228.	JunB-N-25	ATATATTATATTCAATA
229.		
	JunB-N-26	CAAATATATTATATTCAATA
230.	JunB-N-27	TATATTATATTCAAT
231.	JunB-N-28	AATATATTATATTCAAT
		- · · · · · · · · - <del>-</del>
232.	JunB-N-29	TATATTATATTCAA
233.	JunB-N-30	CAAATATATTATATTCAA
234.	JunB-N-31	CAAATATATTATATTCA
235.	JunB-N-32	CAAATATATTATATTC
236.	JunB-N-33	CACAAATATATTATATTC
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237.	JunB-N-34	AAATATATTATATT
238.	JunB-N-35	CAAATATATTATATT
239.	JunB-N-36	CAAATATATTATAT
	·	
240.	JunB-N-37	CACAAATATATTATAT
241.	JunB-N-38	CACAAATATATTAT
242.	JunB-N-39	TACACAAATATATTAT
243.	JunB-N-40	TACACAAATATATTA
244.	JunB-N-41	TAAATACACAAATATATT
245.	JunB-N-42	AATACACAAATATA
246.	JunB-N-43	GTTAAATACACAAATA
247.	JunB-N-44	TGTTAAATACACAA
		ICILIAAA LALALAA
24Ω	<del></del>	
248.	JunB-N-45	TTTAGAGACTAAGT
248. 249.	<del></del>	
249.	JunB-N-45 JunB-N-46	TTTAGAGACTAAGT ATAAACTCTTTAGA
249. 250.	JunB-N-45 JunB-N-46 JunB-N-47	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG
249. 250. 251.	JunB-N-45 JunB-N-46	TTTAGAGACTAAGT ATAAACTCTTTAGA
249. 250.	JunB-N-45 JunB-N-46 JunB-N-47	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG
249. 250. 251. 252.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT
249. 250. 251. 252. 253.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTCTTT
249. 250. 251. 252.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT
249. 250. 251. 252. 253. 254.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTCTTT TAAAAATAAACTC
249. 250. 251. 252. 253. 254. 255.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTCTTT TAAAAAAAA
249. 250. 251. 252. 253. 254. 255.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52 JunB-N-53	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTCTTT CTTAAAATAAACTC TAAAAAGAACAAACA TAAAAAGAACAAACA CAATAAAAAGAACAA
249. 250. 251. 252. 253. 254. 255.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTCTTT TAAAAAAAA
249. 250. 251. 252. 253. 254. 255. 256.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52 JunB-N-53 JunB-N-54	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTC TAAAAAGAACAAACA TAAAAAGAACAAACA CAATAAAAAGAACAA TCAATAAAAAGAACAA
249. 250. 251. 252. 253. 254. 255. 256. 257.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52 JunB-N-53 JunB-N-54 JunB-N-55	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTC TAAAAAGAACAAACA TAAAAAGAACAAACA CAATAAAAAGAACAA TCAATAAAAAGAACAA TCAATAAAAAGAACAA
249. 250. 251. 252. 253. 254. 255. 256. 257. 258. 259.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52 JunB-N-53 JunB-N-54	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTC TAAAAAGAACAAACA TAAAAAGAACAAACA CAATAAAAAGAACAA TCAATAAAAAGAACAA
249. 250. 251. 252. 253. 254. 255. 256. 257.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52 JunB-N-53 JunB-N-54 JunB-N-55	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTC TAAAAAGAACAAACA TAAAAAGAACAAACA CAATAAAAAGAACAA TCAATAAAAAGAACAA TCAATAAAAAGAACAA
249. 250. 251. 252. 253. 254. 255. 256. 257. 258. 259. 260.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52 JunB-N-53 JunB-N-54 JunB-N-55 JunB-N-55	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTC TAAAAAGAACAAACA TAAAAAGAACAAACA CAATAAAAAGAACAA TCAATAAAAAGAACAA TCAATAAAAAGAAC
249. 250. 251. 252. 253. 254. 255. 256. 257. 258. 259.	JunB-N-45 JunB-N-46 JunB-N-47 JunB-N-48 JunB-N-49 JunB-N-50 JunB-N-51 JunB-N-52 JunB-N-53 JunB-N-54 JunB-N-55 JunB-N-55	TTTAGAGACTAAGT ATAAACTCTTTAGA TAAAATAAACTCTTTAG TAAAATAAACTCTTTA TTAAAATAAACTCTTT CTTAAAATAAACTC TAAAAAGAACAAACA TAAAAAGAACAAACA CAATAAAAAGAACAA TCAATAAAAAGAACAA TCAATAAAAAGAAC

261.	JunB-T-1	TGGCGCGGCGGTAGC
	<del>-</del>	
262.	JunB-T-2	GGGCTGGCGGGGGTAG
263.	JunB-T-3	TCGGGGCTGGCGCGGGG
264.	JunB-T-4	TGGGTGCCTGGTCGCGCGTTCTCGGG
	JunB-T-5	AGGGTCCCTGCGGGGCCG
265.		
266.	JunB-T-6	GGGAGGGTCCCTGCGGGG
267.	JunB-T-7	GGGAGGGTCCCTGCGG
		TGGGCCGGGTCCGC
268.	JunB-T-8	
269.	JunB- <b>T-</b> 9	TCCCGGGGTGTAG
270.	JunB-T-10	AGTACTGTCCCGGGGGTGT
271.	JunB-T-11	GGGACACGTTGGGGGGTG
272.	JunB-T-12	GCCGGGGCCCCCGGTAGC
273.	JunB-T-13	CGGGCCCAGCCGGGGC
	<del></del>	CGGGCCCAGCCGGG
274.	JunB-T-14	
275.	Jun <b>B-T-1</b> 5	GGGAGGTGGCTCCGGGCCGG
276.	JunB-T-16	AGGCCGCGCGTGTGGGA
277.	JunB-T-17	GGGTGGCCACCGGCGAAGGG
	_ <del> </del>	
278.	JunB-T-18	AGGGCAGGGACGT
279.	JunB-T-19	TAAAGGGGCAGGGACGT
	JunB-T-20	AGGGGGTGTCCGTAAAGGGG
280.	0mb-1-20	DOCOMANI DOOLO TO
281.	JunD-T-1	GGGGACGCGAACGTGCCGCCG
282.	JunD-T-2	CGGGGAACAAGCGGCCCGGGG
283.	<b>JunD-T-</b> 3	GGCCGTCGGGGGCG
284.	JunD-T-4	GCGGCCGTCGGGGGC
285.	JunD-T-5	AGGGGGTAGGAGGCGGG
286.	JunD-T-6	GCGCTGGGGCCC
287.	JunD-T-7	GGCCGTCGGGGGGT
288.	JunD-T-8	GGGGAGGCCAGCTTC
289.	JunD-T-9	GGCCGCCACCTTGGGG
290.	JunD-T-10	GCGGCCGCCGGGG
291.	JunD-T-11	GGGCGCGCCGCCGGGG
	— ——— — ·	GGGTGGCGGCGG
292.	JunD-T-12	
293.	JunD-T-13	GGGGTGGCGGC
294.	JunD-T-14	TGGGCAGCAGCTGGCAG
	JunD-T-15	CGGGCCCCACGACACC
295.		aaaaaaaaaa maraa ah
295. 296.	JunD-T-16	CGGGGCCCCACGACAC
		GGGGGCCCACGACAC
296.	JunD-T-16	
296. 297.	JunD-T-16 JunD-T-17	GGGCCGCACCCTCTCCAAGTCCGGGG
296.	JunD-T-16 JunD-T-17 ErbB-2-1	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG
296. 297.	JunD-T-16 JunD-T-17	GGGCCGCACCCTCTCCAAGTCCGGGG
296. 297. 298. 299.	JunD-T-16 JunD-T-17 ErbB-2-1 ErbB-2-2	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG
296. 297. 298. 299. 300.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC
296. 297. 298. 299. 300. 301.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC
296. 297. 298. 299. 300.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC
296. 297. 298. 299. 300. 301. 302.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC
296. 297. 298. 299. 300. 301. 302. 303.	JunD-T-16 JunD-T-17 ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC
296. 297. 298. 299. 300. 301. 302. 303. 304.	JunD-T-16 JunD-T-17 ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC
296. 297. 298. 299. 300. 301. 302. 303.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG
296. 297. 298. 299. 300. 301. 302. 303. 304.	JunD-T-16 JunD-T-17 ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-7 ErbB-2-9 ErbB-2-10 ErbB-2-11	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAAACACTTGG GGTAACCTGTGATCTCC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCC CCTGCAGTACTCCC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-14	GCAGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACACTCGG CCCACACATCCCC CCCACACCCCCCCCCC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-14	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311. 312. 313.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-7 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-14 ErbB-2-15 ErbB-2-16	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CCACAAACACTTCCG GCAAACAGTGCCTGGC CGCATCGTGTACTTCCG GCACGTTCCGAGCG
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311. 312. 313. 314.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CGCATCGTGTACTTCCG GCATCGTGTACTTCCG GCACGTTCCGAGCG GCACGTTCCGAGCG
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311. 312. 313.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-12 ErbB-2-13 ErbB-2-14 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CCACAGTGTGTCCTGGC CCACAGTGTGTCCTCCG GCAACCGTGTACTTCCC CCAGTGTGTACTTCCC CCAGTGTGTACTTCCC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311. 312. 313. 314.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CGCATCGTGTACTTCCG GCATCGTGTACTTCCG GCACGTTCCGAGCG GCACGTTCCGAGCG
296. 297.  298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311. 312. 313. 314. 315.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-12 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CCAGAGTCTCGGG GGCATTCACATACTCC CCAGAGTCTCGAGCG CCACGTGTGATCTCCG CCACAGCGTTCCGAGCG CCACGTGCAGCG CCACGTGCAGCG CCAGTGGAGACCTGGG CCTGAGGACACATCAGG
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311. 312. 313. 314. 315. 316. 317.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-15 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19 ErbB-2-20	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAACAGTGCCTGGC CCACAGTGTACTCCC CCAGAGTCTCGAGCG CCCACTGTGTACTCCC CCAGTGTACTCCG CCCATCGTGTACTCCC CCCAGTGTACTCCC CCAGTGGAGCC CCTCACTTGGTTGTGAGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 311. 312. 313. 314. 315. 316. 317. 318.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-13 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19 ErbB-2-20 ErbB-2-21	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CGCATCGTGATCTCCG GCACGTTCCGAGCG CCACGTGCAGCG GCACCTCGAGCG GGTACCAGATACTCC CCAGTGGAGACCTGG CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 309. 310. 311. 312. 313. 314. 315. 316. 317.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-15 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19 ErbB-2-20	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAACAGTGCCTGGC CCACAGTGTACTCCC CCAGAGTCTCGAGCG CCCACTGTGTACTCCC CCAGTGTACTCCG CCCATCGTGTACTCCC CCCAGTGTACTCCC CCAGTGGAGCC CCTCACTTGGTTGTGAGC
296. 297.  298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 311. 312. 313. 314. 315. 315. 316. 317. 318. 319.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-14 ErbB-2-15 ErbB-2-16 ErbB-2-16 ErbB-2-17 ErbB-2-19 ErbB-2-20 ErbB-2-21 ErbB-2-21	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CGCATCGTGATCTCCG GCACGTTCCGAGCG CCACGTGCAGCG GCACCTCGAGCG GGTACCAGATACTCC CCAGTGGAGACCTGG CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 310. 311. 312. 313. 314. 315. 316. 317. 318. 319. 319. 319.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-14 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19 ErbB-2-20 ErbB-2-21 ErbB-2-21 ErbB-2-21	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CCAGAGTCTCGAGCG CCACTGTGTACTTCCG GCACTTCCGAGCG GCACCTTCGAGCG CCTGAGGACACATCAGG CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC GGAAGATGTCCTTCC GCACACTGCTCATGGC GCACACTCCTCATGGC GCACACTGCTCATGGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 310. 311. 312. 313. 314. 315. 316. 317. 318. 319. 320. 321.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-12 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19 ErbB-2-20 ErbB-2-20 ErbB-2-21 ErbB-2-21 ErbB-2-22 ErbB-2-23 ErbB-2-24	GCAGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCCG GCAATCCTGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CCAGTGTGTACTTCCG CCTGCAGTACTCCG GCACTTCCGAGCG GGTACCAGATACTCC CCAGTGGAGACCTGG CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC GGAAGATGTCCTTCC GCACACTGCTCATGGC GCACACTGCTCATGGC GCACACTGCTCATGGC GCACACTGCTCATGGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 310. 311. 312. 313. 314. 315. 316. 317. 318. 319. 319. 319.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-14 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19 ErbB-2-20 ErbB-2-21 ErbB-2-21 ErbB-2-21	GGGCCGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAAACAGTGCCTGGC CCAGAGTCTCGAGCG CCTCCTGGTTCCGGC CCCATCGTGTACTTCC GCAACCTGTGTACTTCC GCAACCTGTGTACTTCC GCACTTCCGAGCG GCTACCAGATACTCC CCAGTGGAGACCTTGG CCTCACTTGGTTGTGAGC GGAAGATGTCCTTCC GCACACTGCTCATGGC GCACACTGCTCATGGC GCACACTCCTCATGGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 309. 310. 311. 312. 313. 314. 315. 316. 317. 318. 319. 320. 321. 322.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-12 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-18 ErbB-2-19 ErbB-2-20 ErbB-2-20 ErbB-2-21 ErbB-2-21 ErbB-2-22 ErbB-2-23 ErbB-2-24	GCAGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTCC CCTGCAGTACTCCG GCAATCCTGGC GCATCCTGGC CCAGAGTCTCAGC CCAGAGTCTCAGC CCTCTGGTTACTCC GCAACCAGTACTCC GCAACCAGTACTCC GCACCTTCGAGCC GCACCTTCCGAGCG CCTCACTGGTTCCC CCAGTGGAGACCTGG CCTCACTTGGTTGTGAGC CCTCACTTGGTTGTGAGC GGAAGATGTCCTTCC GCACACTGCTCATGGC GCACACTGCTCATGGC GCACACTGCTCATGGC
296. 297. 298. 299. 300. 301. 302. 303. 304. 305. 306. 307. 308. 310. 311. 312. 313. 314. 315. 316. 317. 318. 319. 320. 321.	JunD-T-16 JunD-T-17  ErbB-2-1 ErbB-2-2 ErbB-2-3 ErbB-2-4 ErbB-2-5 ErbB-2-6 ErbB-2-7 ErbB-2-8 ErbB-2-9 ErbB-2-10 ErbB-2-11 ErbB-2-12 ErbB-2-13 ErbB-2-14 ErbB-2-15 ErbB-2-15 ErbB-2-16 ErbB-2-17 ErbB-2-19 ErbB-2-20 ErbB-2-20 ErbB-2-21 ErbB-2-21 ErbB-2-22 ErbB-2-23 ErbB-2-24 ErbB-2-25	GCAGCACCCTCTCCAAGTCCGGGG GCAGCAGTCAGTGG CCATTGTCTAGCACGG GGTCTCCATTGTCTAGC GGTGGTATTGTTCAGC GCTGGATCAAGACCC CCACAAAATCGTGTCC CCTTCCACAAAATCGTGTCC GGTTGTTCTTGTGG CCTCTTGGTTGTGC CCAGAGTCTCAAACACTTGG GGTAACCTGTGATCTCTTCC CCTGCAGTACTCGG GGCATTCACATACTCC GCAACAGTGCCTGGC CGCATCGTGTACTTCCG GCACTTCCGAGCG CCTCTGGGTTCCGAGCG GGTACCAGATACTCC CCAGTGGAGACCTTGG CCTCACTTGGTTGTGC CCTGAGGACACTCCC CCAGTGGAGACCTTGG CCTCACTTGGTTGTGAGC GCACACTCCTTCC CCACACATCACTCC CCACACATCACCC CCACACATCACCC

Fig: 3 - 5

324.	ErbB-2-27	CCTTCTGGTTCACACTGG
325.	ErbB-2-28	CATGGTGCTCACTGCG
326.	ErbB-2-29	CTTGGTTGTGAGCG
327.	ErbB-2-30	
		GGACAGGCAGTCAC
328.	ErbB-2-31	GTCACCTCTTGGTTGTGC
329.	ErbB-2-32	CCAGAGTCTCAAACAC
330.	ErbB-2-33	CACATACTCCCTGG
331.	ErbB-2-34	GACCAGCACGTTCCG
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332.	ErbB-2-35	GTTGGTGTCTATCAGTG
333.	ErbB-2-36	CCCTGGTAGAGGTG
334.		
	ErbB-2-37	CTCAAACACTTGGAGC
335.	ErbB-2-38	CACACATCACTCTGGTGG
336.		
	ErbB-2-39	GCACAGACAGTGCGC
337.	ErbB-2-40	CATGGCAGCAGTCAG
338.	ErbB-2-41	CTGCTCATGGCAGCAG
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339.	ErbB-2-42	CATCTGGAAACTTCCAGATG
340.	ErbB-2-43	CTGGAAACTTCCAG
341.	ErbB-2-44	CATAACTCCACACATCACTC
342.	ErbB-2-45	CACCATAACTCCACACATC
343.	ErbB-2-46	CTGGTGGGTGAACC
344.	ErbB-2-47	CGGATTACTTGCAGG
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345.	ErbB-2-48	CGCTAGGTGTCAGCG
346.	ErbB-2-49	GCCATCACGTATGC
347.		
	ErbB-2-50	GCATACACCAGTTCAGC
348.	ErbB-2-51	CCATCAAATACATCGG
349.	ErbB-2-52	
	_	CCAGCAGAAGTCAGG
350.	ErbB-2-53	GCTTCATGTCTGTGC
351.	ErbB-2-54	GGTGAGTTCCAGGTTTCC
352.	ErbB-2-55	CCACAAAATCGTGTCCTGG
353.	ErbB-2-56	CCCTTACACATCGG
354.	ErbB-2-57	GCAGCTCACAGATGC
355.	ErbB-2-58	GCACTGGTAACTGC
356.		
	ErbB-2-59	CCTGGATATTGGCACTGG
357 <i>.</i>	ErbB-2-60	CCAGCAAACTCCTGG
358.	ErbB-2-61	
		GCAGAAATGCCAGGC
359.	ErbB-2-62	CCATTGTGCAGAATTCG
360.	ErbB-2-63	CCCTGCAGTACTCGG
361.	ErbB-2-64	GGCATTCACATACTCCC
362.	ErbB-2-65	GGTCAGGTTTCACACC
363.	ErbB-2-66	CCAGGTCCACACAGG
364.	ErbB-2-67	CCTTGTCATCCAGG
365.		
	ErbB-2-68	GGATCCCAAAGACC
366.	ErbB-2-69	CCTCAACACTTTGATGG
367.	ErbB-2-70	
		GCTGTGTCACCAGC
368.	ErbB-2-71	GGTCTAAGAGGCAGCC
369.	ErbB-2-72	GGCAATCTGCATACACC
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370.	ErbB-2-73	CCTGTGTACGAGCC
371.	ErbB-2-74	CCATCCACTTGATGG
372.		· ·
	ErbB-2-75	CCCACACAGTCACACC
373.	ErbB-2-76	CCATCGTAAGGTTTGG
374.	ErbB-2-77	CCTTTTCCAGCAGG
		·
3 <b>75.</b>	ErbB-2-78	GGAGAATTCAGACACC
376.	ErbB-2-79	CCAAGTCCTCATTCTGG
377.	ErbB-2-80	CCATCAGTCTCAGAGG
378.	ErbB-2-81	CCTTTGAAGGTGCTGG
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379.	ErbB-2-82	GGCATGGCAGGTTCC
380.	ErbB-2-83	CCTGGCATGGCAGG
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381.	ErbB-2-N-1	AGATGTATAGGTAA
382.	ErbB-2-N-2	ATTTTCACATTCTC
383.	ErbB-2-N-3	AATTTTCACATTCTC
384.	ErbB-2-N-4	
		AATTTTCACATTCT
385.	ErbB-2-N-5	GAATTTTCACATTC
386.	ErbB-2-N-6	GGAATTTTCACATT
387.	ErbB-2-N-7	AGATTTCTTTGTTG
388.	ErbB-2-N-8	AAGATTTCTTTGTTG
389,	ErbB-2-N-9	AAGATTTCTTTGTT
Fig. 2	·	
Fig. 3 - 6		

390.	ErbB-2-N-10	TAAGATTTCTTTGTT
391.	ErbB-2-N-11	CTAAGATTTCTTTGTT
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392.	ErbB-2-N-12	TAAGATTTCTTTGT
393.	ErbB-2-N-13	CTAAGATTTCTTTGT
394.	ErbB-2-N-14	CTAAGATTTCTTTG
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395.	ErbB-2-N-15	TCTAAGATTTCTTT
396.	ErbB-2-N-16	GTCTAAGATTTCTTT
397.	ErbB-2-N-17	GTCTAAGATTTCTT
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398.	ErbB-2-N-18	TTCGTCTAAGATTT
399.	ErbB-2-N-19	ATTTTGACATGGTT
400.	ErbB-2-N-20	AATTTTGACATGGTT
401.	ErbB-2-N-21	AATTTTGACATGGT
402.	ErbB-2-N-21	TAATTTTGACATGGT
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403.	ErbB-2-N-23	TAATTTTGACATGG
404.	ErbB-2-N-24	GTAATTTTGACATG
405.	ErbB-2-N-25	TGTAATTTTGACATG
406.	ErbB-2-N-26	TGTAATTTTGACAT
407.	ErbB-2-N-27	TCTGTAATTTTGACAT
408.	ErbB-2-N-28	CTGTAATTTTGACA
409.	ErbB-2-N-29	TCTGTAATTTTGACA
410.	ErbB-2-N-30	TCTGTAATTTTGAC
411.	ErbB-2-N-31	GTCTGTAATTTTGA
412.	ErbB-2-N-32	AAGTCTGTAATTTTGA
413.	ErbB-2-N-33	AGTCTGTAATTTTG
414.	ErbB-2-N-34	AAGTCTGTAATTTTG
415.	ErbB-2-N-35	AAGTCTGTAATTTT
416.	ErbB-2-N-36	GAAGTCTGTAATTTT
417.	ErbB-2-N-37	GAAGTCTGTAATTT
418.	ErbB-2-N-38	ATGTAGACATCAAT
419.	ErbB-2-N-39	ATCATCCAACATTT
420.	ErbB-2-N-40	AATCATCCAACATTT
421.	ErbB-2-N-41	AATCATCCAACATT
422.	ErbB-2-N-42	ACCATCAAATACAT
423.	ErbB-2-N-43	AAAAACGTCTTTGA
424.	ErbB-2-N-44	TTTTGTTCTTAGACA
425.	ErbB-2-N-45	TTTTGTTCTTAGAC
426.	ErbB-2-N-46	TAAACAGAAAAGCA
427.	ErbB-2-N <b>-</b> 47	ACTAAACAGAAAAG
428.	ErbB-2-N-48	AAACTAAACAGAAAAG
429.	ErbB-2-N-49	AACTAAACAGAAAA
430.	ErbB-2-N-50	AAACTAAACAGAAAA
431.	ErbB-2-N-51	AAACTAAACAGAAA
432.	ErbB-2-N-52	TAAAAACTAAACAGAAA
433.	ErbB-2-N-53	AAAACTAAACAGAA
434.	ErbB-2-N-54	GTAAAAACTAAACAGAA
435.	ErbB-2-N-55	AAAAACTAAACAGA
436.	ErbB-2 <b>-</b> N <b>-</b> 56	TAAAAACTAAACAGA
437.	ErbB-2-N-57	TAAAAACTAAACAG
438.	ErbB-2-N-58	
439.	ErbB-2-N-59	AAAAAGTAAAACTAAACA
440.	ErbB-2-N-60	AGTAAAAACTAAAC
441.	ErbB-2-N-61	AAAAAAGTAAAAACTAAAC
		<del></del>
442.	ErbB-2-N-62	AAGTAAAACTAAA
443.	ErbB-2-N-63	AAAAAAGTAAAACTAAA
444.	ErbB-2-N-64	AAAGTAAAACTAA
445.	ErbB-2-N-65	AAAAGTAAAAACTA
446.	ErbB-2-N-66	AAAAAAGTAAAAACTA
447.	ErbB-2-N-67	AAAAGTAAAACT
448.	ErbB-2-N-68	AAAAAAGTAAAAACT
449.	ErbB-2-N-69	AAAAAAGTAAAAAC
		CAAAAAAGTAAAAAC
450.	ErbB-2-N-70	
451.	ErbB-2-N-71	AAAAAAGTAAAAA
452.	ErbB-2-N-72	CAAAAAAGTAAAA
453.	ErbB-2-N-73	
454.	ErbB-2-N-74	AAACAAAAAAGTA
455.	ErbB-2-N-75	CAAAACAAAAAAGTA
456.	ErbB-2-N-76	CAAAACAAAAAAGT
T - ~	2 7	
Fig.	3 - 7	

457.		ErbB-2-N-77	CAAAACAAAAAAG
458.		ErbB-2-N-78	CTTTAAAAAAACAAAAC
459.		ErbB-2-N-79	TCTTTAAAAAAACAAA
460.			•
		ErbB-2-N-80	GTCTTTAAAAAAACAAA
461.		ErbB-2-N-81	GTCTTTAAAAAAAACA
462.		ErbB-2-N-82	GTCTTTAAAAAAAC
463.		ErbB-2-N-83	TTTATTTCGTCTTT
464.		ErbB-2-N-84	TCTTTATTTCGTCT
465.		ErbB-2-N-85	TATTTGCAAATGGA
466.		ErbB-2-N-86	TATATTTGCAAATGG
467.		ErbB-2-N-87	TATATTTGCAAATG
468.		ErbB-2-N-88	CAAAATATATTTGCAAATG
469.		ErbB-2-N-89	
		<del></del>	CAAAATATATTTGCAAAT
470.		ErbB-2-N-90	CAAAATATATTTGCA
471.		ErbB-2-N-91	CAAAATATATTTGC
472.		ErbB-2-N-92	TTCCAAAATATATTTG
473.		ErbB-2-N-93	TTTTCCAAAATATATTT
474.		ErbB-2-N-94	
			GTTTTCCAAAATATATT
475.		ErbB-2-N-95	GTTTTCCAAAATAT
476.		c-fos-1	GGTTAGGCAAAGCC
477.		c-fos-2	CCGAGAACATCATCGTGG
478.		c-fos-3	CCGAGAACATCATCGTG
479.		c-fos-4	
			CCGAGAACATCATCG
480.		c-fos-5	CGTAGTCTGCGTTGAAGC
481.		c-fos-6	CCATGCTGGAGAAGG
482.		c-fos-7	CCGTGCAGAAGTCC
483.		c-fos-8	GGAATGAAGTTGGC
		_	
484.		c-fos-8	TGACCGTGGGAATG
485.		c-fos-10	TGGCAGTGACCGTG
486.		c-fos-11	AGATGGCAGTGACC
487.		c-fos-12	CGAGATGGCAGTGACC
488.	•		
		c-fos-13	CCAGCCACTGCAGG
489.		c-fos-14	GCACCAGCCACTGC
490.		c-fos-15	CCCTGGAGTAAGCC
491.		c-fos-16	GGAGATAACTGTTCCACC
492.		c-fos-17	
		_	GGAGATAACTGTTCC
493.		c-fos-18	CTTCTAGTTGGTCTG
494.		c-fos-19	CATCTTCTAGTTGG
495.		c-fos-20	TCTCATCTTCTAGTTGG
496.		c-fos-21	CTGCAAAGCAGACTTCTC
497.		c-fos-22	
			CCTTCAGCAGGTTGG
498.		c-fos-23	CCCAGGTCATCAGG
499.		c-fos-24	CCAGTCAGATCAAGG
500.		c-fos-25	GGTGAAGGCCTCCTC
501.		c-fos-26	CAGGGTGAAGGCCTC
502.		c-fos-27	CCTGGATGATGCTGG
503.		c-fos-28	CCACTGTGCAGAGG
504.		c-fos-29	GGAGTACAGGTGACC
505.		c-fos-30	GCTCATTGCTGCTGC
506.		c-fos-31	GGAAGGCTCATTGCTGC
		3 232 22	
507.		c-fos-N-1	TTTTCTCTTCTT
		_	-
508.		c-fos-N-2	ATCTTATTCCTTTC
509.		c-fos-N-3	CATCTTATTCCTTT
510.		c-fos-N-4	TAGTTTTTCCTTCT
511.		c-fos-N-5	TCTAGTTTTTCCTT
512.			
		c-fos-N-6	AACTCTAGTTTTTC
513.		c-fos-N-7	GAACTCTAGTTTTT
514.		c-fos-N-8	TGAACTCTAGTTTTT
515.		c-fos-N-9	ATGAACTCTAGTTTTT
516.		c-fos-N-10	
			TGAACTCTAGTTTT
517.		c-fos-N-11	ATGAACTCTAGTTTT
518.		c-fos-N-12	ATGAACTCTAGTTT
519.		TGF-82-1	GCACACAGTAGTGC
Fig.	3 - 8		

		11 / 30
520.	TGF-\$2-2	GCAGGATCAGAAAAGC
521.	TGF-\$2-3	GCAGGTAGACAGGC
522.	TGF-\$2-4	GCTTGCTCAGGATCTGC
523.	TGF-B2-5	GCAAGTCCCTGGTGC
524.		
	TGF-ß2-6	CCTGGAGCAAGTCC
525.	TGF-\$2-7	CGTAGTACTCTTCGTCG
526.	TGF-\$2-8	CGTAGTACTCTTCG
527.	TGF-ß2-9	GTAAACCTCCTTGG
528.	TGF-ß2-10	GTCTATTTTGTAAACCTCC
529.	TGF-£2-11	GCATGTCTATTTTGTAAACC
530.	TGF-£2-12	GGCATCAAGGTACCC
531.	TGF-ß2-13	GGCATCAAGGTACC
532.	TGF-ß2-14	GCTTTCACCAAATTGGAAGC
533.	TGF-\$2-15	GAGAATCTGATATAGCTC
534.	TGF-ß2-16	GGAGATGTTAAATCTTTGG
535.	TGF-B2-17	GCTGTCGATGTAGC
536.		
	TGF-82-18	CCAGGTTCCTGTCTTTATGG
537.	TGF-B2-19	CAGCAGGGACAGTG
538.	TGF-ß2-20	CTTGCTTCTAGTTCTTCAC
539.	TGF-ß2-21	GCCATCAATACCTGC
540.	TGF-£2-22	GGTGCCATCAATACC
<b>541.</b>	TGF-ß2-23	CCACTGGTATATGTGG
542.	TGF-ß2-24	GGACTTTATAGTTTTCTG
543.	TGF-132-25	CTCAAGTCTGTAGGAG
544.	TGF-B2-26	GGTCTGTTGTGACTC
545.	TGF-ß2-27	
·		CAATTATCCTGCACATTTC
546.	TGF-52-28	GCAGCAATTATCCTGC
547.	TGF-B2-29	GGCAGCAATTATCC
548.	TGF-\$2-30	GGTTCGTGTATCCATTTCC
549.	TGF-ß2-31	GCACAGAAGTTGGC
550.	TGF-£2-32	CCAGCACAGAAGTTGG
551.	TGF-\$2-33	GTGCTGAGTGTCTG
552.	TGF-\$2-34	CCTGCTGTGCTGAGTG
553.	TGF-ß2-35	GCTCAGGACCCTGC
554.	TGF-\$2-36	GCAGCAAGGAGAAGC
555.	TGF-\$2-37	CCAATGTAGTAGAGAATGG
556.	TGF-B2-37	
336.	1GF-152-36	GCTGCATTTGCAAG
557.	TGF-B2-N-1	AAAAAAGAAATCAA
558.	TGF-\$2-N-2	AAAAAAAGAAATCAA
559.	TGF-£2-N-3	AAAAAAAGAAATCAA
560.	TGF-\$2-N-4	TAAAAAAAGAAATCAA
561.	TGF-\$2-N-5	
562.	TGF-\$2-N-6	AATAAAAAAAGAAATCAA
563.	TGF-\$2-N-7	GAATAAAAAAAGAAAT
564.	TGF-B2-N-8	
565.	TGF-B2-N-9	
566.	<del>-</del>	
	TGF-\$2-N-10	TCAGAATAAAAAA
567.	TGF-82-N-11	TTGTTTTTAAAAGT
568.	TGF-£2-N-12	AGTTGTTTTAAAA
569.	TGF-ß2-N-13	AAGTTGTTTTAAAA
570.	TGF-B2-N-14	AAAGTTGTTTTAAAA
571.	TGF-\$2-N-15	AAAAGTTGTTTTAAAA
572.	TGF-B2-N-16	AAAAAGTTGTTTTAAAA
573.	TGF-\$2-N-17	AAAAAGTTGTTTTAAAA
574.	TGF-B2-N-18	AAAAAAAGTTGTTTTAAAA
575.	TGF-B2-N-19	
	<del>-</del>	AAAAAAAAGTTGTTTTAAA
576.	TGF-£2-N-20	TTTTTAAAAAAGTG
5 <b>77.</b>	TGF-\$2-N-21	TTTTTTAAAAAGTG
578.	TGF-ß2-N-22	ATTTTTTAAAAAAGTG
579.	TGF-£2-N-23	- CATTTTTTAAAAAAGT
580.	TGF~ß2-N-24	GCATTTTTAAAAAA
581.	TGF-\$2-N-25	TGCATTTTTAAAAAA
582.	TGF-ß2-N-26	AGCTTATTTTAAAT
583.	TGF-\$2-N-27	AAGCTTATTTTAAAT
584.	TGF-ß2-N-28	TAAGCTTATTTTAAAT
585	TGF-82-N-29	TGTAATTATTAGAT
Fig.	3 ~ 9	

586. 587. 588. 589. 590. 591. 592. 593. 594. 595.		TGF-\$2-N-30 TGF-\$2-N-31 TGF-\$2-N-32 TGF-\$2-N-33 TGF-\$2-N-34 TGF-\$2-N-35 TGF-\$2-N-35 TGF-\$2-N-36 TGF-\$2-N-37 TGF-\$2-N-37 TGF-\$2-N-39 TGF-\$2-N-40	ATGTAATTATTAGAT TGATGTAATTATTA ATGATGTAATTATTA ATGGTATTATATAA TATGGTATTATATAA TTATGGTATTATATAA TTTATGGTATTATATAA ATTTATGGTATTATATAA ATTTATGGTATTATATATA
5990123456789012345666666666666666666666666666666666666		rb-1 rb-2 rb-3 rb-4 rb-5 rb-6 rb-7 rb-8 rb-9 rb-10 rb-11 rb-12 rb-13 rb-14 rb-15 rb-16 rb-17 rb-18 rb-19 rb-20 rb-21 rb-22 rb-23 rb-24 rb-25 rb-26 rb-27 rb-28 rb-29 rb-30 rb-31 rb-32 rb-33 rb-34 rb-35 rb-36 rb-37 rb-38 rb-37 rb-38 rb-40 rb-41 rb-42 rb-43 rb-44	GGCATGACGCCTTTC GCATGACGACGTTTC GCCTGACGAGAGGC CTCAAGCCTGACGAG CCACAGTTCCTTTTTC GCTGCAATAAAGATACAG GCTGCAATAAAGATACAG GCTGCAATAAAGATAC GGACACTGATTCTATG GCATTATCAACTTTGG ACTTTTAGCACCAATG CCAAGAAACTTTTAGCACC CCAGATCATCTTCC AGTCAAGGACACATAG TCTTTGAGCAACATAG TCTTTGAGCAACATAG GGGTATAACAGCTG GAGGTGAACCATTAATGG TCTTCGTATCGTTTAG GCATTCATTCCTG GGATTCACTTCC GCAATAAAGACATTCTC GCAATAAAGACATTCTC GCCAATAAAGACATTCTG CCAGAATCAAGATTCTG CCAGAATCAAGATTCTG CCAGAATCAAGATTCTG GACAAAACTTGTTCCAGAATC GGAAAGACAAATCTGTTCC GATTAAGAGGACAAGC GGAAGATTAACAGG GCAGTGTGATTATTCTGG GCAGTTTACAGG GCATTTGCAGTAGAATTTAC CAGTGAAAGACAATTTTCTG GGAGATCTTACAGG GCATTTTTTTTTT
641. 642. 643. 644. 645. 646. 647. 648. 649. 650. Fig.	3 - 10	rb-45 rb-N-1 rb-N-2 rb-N-3 rb-N-4 rb-N-5 rb-N-6 rb-N-7 rb-N-8 rb-N-9	GGTGTACACAGTGTCC TATCTTTAATTTCT TCTTTTGAATATAA TTCTTTTGAATATAA TTTCTTTTGAATATAA TTTTCTTTTTGAATATAA TTTTCTTTTTGAATATAA ATTTCTTTTTTTTTT

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SS2.	651.	rb-N-10	AGTTAAAGAATTTAT
C63.   rb-N-12			<del></del>
654.   hb-N-14   TITAGITAGITAAA     655.   hb-N-14   ATTICTITIAGITAA     656.   hb-N-15   ATTICTITIAGITAA     657.   hb-N-16   AATTICTITIAGITAA     658.   hb-N-17   AICAATICTITIA     659.   hb-N-18   ATTICTITIAGITAA     659.   hb-N-18   ATTICTITITA     660.   hb-N-19   AATATICTITITA     660.   hb-N-19   AATATICTITITA     661.   hb-N-20   AAATATIAGITCA     662.   hb-N-21   CAAATATIAGITCA     663.   hb-N-22   TCAAATATIAGITCA     664.   hb-N-23   TCTCAAATATIATAA     665.   hb-N-24   AATTINITICAGITA     666.   hb-N-26   TAATAAAATATIATAA     667.   hb-N-26   TAATAAAAATATIATAA     668.   hb-N-26   TAATAAAAATATIATAA     669.   hb-N-28   HTGCLTAATAAAATA     671.   hb-N-38   HTGCLTAATAAAATA     672.   hb-N-31   TAATAAAAATAGICAA     673.   hb-N-32   TITAATAAAATAGICAA     674.   hb-N-33   TITAATAAAATAGICAA     675.   hb-N-34   GITTAATAAAATAGICAA     676.   hb-N-36   GAGTITAATAAATAGI     677.   hb-N-36   GAGTITAATAAATAGI     678.   hb-N-37   AGAGTITAATAAATAGI     679.   hb-N-38   AATAATICTICTATA     680.   hb-N-39   TATATITAAAATAGI     681.   hb-N-40   AICATATATAAATAGI     682.   hb-N-41   AATAAACATITITICA     683.   hb-N-42   AATAAACATITITICA     684.   hb-N-43   AATAATATICTICATA     685.   hb-N-44   AATAAACATITITICA     686.   hb-N-45   TATATITAACATTITITICA     686.   hb-N-46   TITAGAATAACATITITICA     686.   hb-N-47   TATATITAACATTITITICA     686.   hb-N-48   AATAAACATITITICA     687.   hb-N-59   AATAATACATITITICA     688.   hb-N-69   AATAAACATITITICA     689.   hb-N-69   AATAAACATITITICA     699.   hb-N-69   AATAAACATITITICA     699.   hb-N-69   AATAAACATITITICA     699.   hb-N-60   AAGAGTAAAACATITITICA     699.   hb-N-61   AAAGAGTAAAACATITITICA     700.   hb-N-66   AAAGAGTAAAATATITICA     701.   hb-N-60   AAAGAGTAAAAATAT			
656.			TTTAGTAAGTTAAA
657.	655.	rb-N-14	TTTTAGTAAGTTAAA
658.	656.	rb-N-15	ATTTCTTTTAGTAA
659.			
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663.			
664.			
665.			
666.			
668.			
669.	667.	rb-N-26	TAATAAAAATGTGAT
670.	668.	rb-N-27	TAGCTAATAAAAT
671.	669.	rb-N-28	TTAGCTAATAAAAAT
672.			
673.			
674.			
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679.			
680. rb-N-39 TATATTACATTCAT 681. rb-N-40 ATCTATATTACATT 682. rb-N-41 ATAAACATTTTCA 683. rb-N-42 AATAACATTTTCA 684. rb-N-43 AATAACATTTTCA 685. rb-N-44 GAAATAACATTTTT 686. rb-N-45 TGAAATAACATTTTT 686. rb-N-45 TGAAATAACATTTTT 687. rb-N-46 TTGAAATAACATTTTT 688. rb-N-47 TTTGAAATAACATTTTT 689. rb-N-48 TTTTGAAATAACATTTTT 690. rb-N-49 TTTTGAAATAACATTTTT 691. rb-N-50 ATTTTTGAAATAACATTTT 692. rb-N-51 AATTTTGAAATAAACATTTT 693. rb-N-52 AAATTTTGAAATAAACATTTT 694. rb-N-53 AAATTTTGAAATAAACATT 695. rb-N-54 TAAAATTTTTGAAATAACATT 696. rb-N-55 ATAAAATTTTTGAAATAACAT 697. rb-N-56 TATAAAATTTTTGAAATAACAT 698. rb-N-57 GTATAAAATTTTTGAAATAAC 699. rb-N-58 GGTATAAAATTTTT 700. rb-N-58 GGTATAAAATTTTT 701. rb-N-60 AAGGTATAAAATTTTT 702. rb-N-60 AAGGTATAAAATTTTT 703. rb-N-61 AAAGGTATAAAATTTT 704. rb-N-62 AAAAGGTATAAAATTTTT 705. rb-N-64 ATAAAGGTATAAAATTTTT 706. rb-N-65 TTTAGAAAGTTTTT 707. rb-N-66 AAAGGTATAAAATTTTT 708. rb-N-67 TAAAAGGTATAAAATTTTT 709. rb-N-68 TTAAAAGGTATAAAATTTTT 709. rb-N-68 TTAAAAGGTATAAAATTTTT 709. rb-N-68 TTAAAAGGTATAAAATTTTT 709. rb-N-69 TTAAAAGGTATAAAATTTTT 710. rb-N-69 TTAAAGAGTAATAATTCTT 711. rb-N-70 TTTTAAGATAAATTTCTT 712. rb-N-71 TTTTTAAGATAAATTTCTT 713. rb-N-72 ATTTTTAAGATAAATTTCTT 714. rb-N-73 TATTTTAAGATAAATT			
682. rb-N-41 ATAAACATTTTCA 683. rb-N-42 AATAAACATTTTCA 684. rb-N-43 AAATAAACATTTTCA 685. rb-N-44 GAAATAAACATTTTT 686. rb-N-45 TGAAATAAACATTTTT 687. rb-N-46 TTGAAATAAACATTTTT 688. rb-N-47 TTTTGAAATAACATTTTT 689. rb-N-48 TTTTGAAATAAACATTTTT 690. rb-N-49 TTTTTGAAATAAACATTTTT 691. rb-N-50 AATTTTGAAATAAACATTTT 692. rb-N-51 AAATTTTTGAAATAAACATTTT 693. rb-N-52 AAATTTTTGAAATAAACATTT 694. rb-N-53 AAAATTTTTGAAATAAACATT 695. rb-N-54 TAAAATTTTGAAATAAACAT 696. rb-N-55 ATAAAATTTTGAAATAAACAT 697. rb-N-56 TATAAAATTTTTGAAATAACA 698. rb-N-57 GTATAAAATTTTTGAAATAAAC 699. rb-N-58 GGTATAAAATTTTT 700. rb-N-59 AGGTATAAAATTTT 701. rb-N-60 AGGTATAAAATTTT 702. rb-N-61 AAAGGTATAAAATTTT 703. rb-N-62 AAAGGTATAAAATTTT 704. rb-N-62 AAAGGTATAAAATTTTT 705. rb-N-62 AAAGGTATAAAATTTTT 706. rb-N-63 TAAAAGGTATAAAATTTTT 707. rb-N-64 ATAAAGGTATAAAATTTTT 708. rb-N-65 TTTAAGAGTATAAATTTTT 709. rb-N-66 AAGGTATAAAATTTTT 709. rb-N-67 TAAAAGGTATAAAATTTTT 709. rb-N-68 TTTAAGAGTATAAATTTTT 710. rb-N-69 TTTAAGATAAATTTTT 711. rb-N-70 TTTTAAGATAAATTTCTT 712. rb-N-71 TTTTAAGATAAATTTCTT 713. rb-N-72 ATTTTTAAGATAAATTTCTT 714. rb-N-73 TATTTTTAAGATAAATTTCT 715. rb-N-74 TTTTTTTTAAGATAAATTT			TATATTACATTCAT
683.	681.	rb-N-40	ATCTATATTACATT
684. rb-N-43 AAATAACATTITCA 685. rb-N-44 GAAATAACATTITT 686. rb-N-45 TGAAATAACATTITT 687. rb-N-46 TTGAAATAACATTITT 688. rb-N-47 TTTGAAATAACATTITT 689. rb-N-48 TTTTGAAATAACATTITT 690. rb-N-49 TTTTGAAATAACATTITT 691. rb-N-50 ATTTTTGAAATAACATTITT 692. rb-N-51 AAATTTTGAAATAACATTITT 693. rb-N-52 AAATTTTTGAAATAACATT 694. rb-N-53 AAAATTTTTGAAATAACATT 695. rb-N-54 TAAAATTTTTGAAATAACAT 696. rb-N-55 ATAAAATTTTTGAAATAACA 696. rb-N-56 TATAAAATTTTTGAAATAAC 697. rb-N-56 TATAAAATTTTTGAAATAAC 698. rb-N-57 GTATAAAATTTTTAAAT 699. rb-N-58 GGTATAAAATTTTT 700. rb-N-59 AGGTATAAAATTTT 701. rb-N-60 AAAGGTATAAAATTTT 702. rb-N-61 AAAGGTATAAAATTTT 703. rb-N-62 AAAAGGTATAAAATTTTT 704. rb-N-63 TAAAAGGTATAAAATTTTT 705. rb-N-64 ATAAAAGGTATAAAATTTTT 706. rb-N-65 TTTAGAAGATTTTT 707. rb-N-66 AAGATAAATTTTT 708. rb-N-67 TAAAAAGTTTTTT 710. rb-N-69 TTTAAGATAAAATTTTT 711. rb-N-70 TTTTAAGATAAATTTTT 712. rb-N-71 TTTTTAAGATAAATTTTT 713. rb-N-72 ATTTTTAAGATAAATTTTT 714. rb-N-73 TAATTTTTAAGATAAATTTTT 715. rb-N-76 CTTTATTTTAAGATAAATT		rb-N-41	ATAAACATTTTCA
685. rb-N-44 GRAATAACATTTTT 686. rb-N-45 TGRAATAACATTTTT 687. rb-N-46 TTGRAATAACATTTTT 688. rb-N-47 TTTGAAATAACATTTTT 689. rb-N-48 TTTTGAAATAACATTTTT 690. rb-N-49 TTTTGAAATAACATTTTT 691. rb-N-50 ATTTTTGAAATAACATTTTT 692. rb-N-51 AATTTTTGAAATAACATTTT 693. rb-N-52 AAATTTTTGAAATAACATTTT 694. rb-N-53 AAAATTTTTGAAATAACATT 695. rb-N-54 TAAAATTTTGAAATAACAT 695. rb-N-55 ATAAAATTTTGAAATAACAT 696. rb-N-56 TATAAAATTTTTGAAATAACA 697. rb-N-56 TATAAAATTTTTGAAATAACA 698. rb-N-57 GTATAAAATTTTT 699. rb-N-58 GGTATAAAATTTT 700. rb-N-59 AGGTATAAAATTTT 701. rb-N-60 AGGTATAAAATTTT 702. rb-N-61 AAAGGTATAAAATTTT 703. rb-N-62 AAAGGTATAAAATTTT 704. rb-N-63 TAAAAGTATAAATTTT 705. rb-N-64 ATAAAGGTATAAAATTTT 706. rb-N-65 TTTAGAATAAATTTT 707. rb-N-66 AAGGTATAAAATTTT 708. rb-N-65 TTTAGAATAATTTT 709. rb-N-66 AAGATAAATTTTT 710. rb-N-69 TTTAGAATAAATTTT 711. rb-N-70 TTTTAAGATAAATTTTT 712. rb-N-71 TTTTAAGATAAATTTTT 713. rb-N-72 ATTTTAAGATAAATTTTT 714. rb-N-73 TATTTTAAGATAAATT			
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706.       rb-N-65       TTTAGAAAGATTT         707.       rb-N-66       AAGATAAATTTCTT         708.       rb-N-67       TAAGATAAATTTCTT         709.       rb-N-68       TTTAAGATAAATTTCTT         710.       rb-N-69       TTTTAAGATAAATTTCTT         711.       rb-N-70       TTTTTAAGATAAATTTCTT         712.       rb-N-71       TTTTTAAGATAAATTTCTT         713.       rb-N-72       ATTTTTAAGATAAATTTCTT         714.       rb-N-73       TATTTTTAAGATAAATTTCT         715.       rb-N-74       TTATTTTTAAGATAAATT         716.       rb-N-75       TTTATTTTAAGATAAATT         717.       rb-N-76       CTTTATTTTAAGATAAAT	704.	rb-N-63	TAAAAGGTATAAAATTTTT
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708.       rb-N-67       TAAGATAAATTTCTT         709.       rb-N-68       TTAAGATAAATTTCTT         710.       rb-N-69       TTTAAGATAAATTTCTT         711.       rb-N-70       TTTTAAGATAAATTTCTT         712.       rb-N-71       TTTTTAAGATAAATTTCTT         713.       rb-N-72       ATTTTTAAGATAAATTTCTT         714.       rb-N-73       TATTTTTAAGATAAATTTCT         715.       rb-N-74       TTATTTTTAAGATAAATT         716.       rb-N-75       TTTATTTTAAGATAAATT         717.       rb-N-76       CTTTATTTTAAGATAAAT			
709. rb-N-68 TTAAGATAAATTTCTT 710. rb-N-69 TTTAAGATAAATTTCTT 711. rb-N-70 TTTTAAGATAAATTTCTT 712. rb-N-71 TTTTAAGATAAATTTCTT 713. rb-N-72 ATTTTAAGATAAATTTCTT 714. rb-N-73 TATTTTAAGATAAATTTCTT 715. rb-N-74 TTATTTTAAGATAAATT 716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAAT			T
710. rb-N-69 TTTAAGATAAATTTCTT 711. rb-N-70 TTTTAAGATAAATTTCTT 712. rb-N-71 TTTTAAGATAAATTTCTT 713. rb-N-72 ATTTTAAGATAAATTTCTT 714. rb-N-73 TATTTTAAGATAAATTTCT 715. rb-N-74 TTATTTTAAGATAAATT 716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAAT			
711. rb-N-70 TTTTAAGATAAATTTCTT 712. rb-N-71 TTTTAAGATAAATTTCTT 713. rb-N-72 ATTTTTAAGATAAATTTCTT 714. rb-N-73 TATTTTTAAGATAAATTTCT 715. rb-N-74 TTATTTTAAGATAAATT 716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAAT			
712. rb-N-71 TTTTAAGATAAATTTCTT 713. rb-N-72 ATTTTAAGATAAATTTCTT 714. rb-N-73 TATTTTAAGATAAATTTCT 715. rb-N-74 TTATTTTAAGATAAATT 716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAAT			
713. rb-N-72 ATTTTAAGATAAATTTCTT 714. rb-N-73 TATTTTAAGATAAATTTCT 715. rb-N-74 TTATTTTAAGATAAATT 716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAAT			
714. rb-N-73 TATTTTAAGATAAATTTCT 715. rb-N-74 TTATTTTAAGATAAATT 716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAATT			
715. rb-N-74 TTATTTTAAGATAAATT 716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAATT			
716. rb-N-75 TTTATTTTAAGATAAATT 717. rb-N-76 CTTTATTTTAAGATAAAT			
	716.		TTTATTTTTAAGATAAATT
Fig. 3 - 11	717.	rb-N-76	CTTTATTTTTAAGATAAAT
	Fia 3 - 1	1	
	* ± ± ± · · · · · · · · · · · · · · · ·	<del>-</del> -	
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710		wh N 77	
718.		rb-N-77	TCTTTATTTTTAAGATAAAT
719.		rb-N-78	ATCTTTATTTTTAAGATAAA
720.		rb-N-79	ATCTTTATTTTAA
721.		rb-N-80	GATCTTTATTTTAA
722.		rb-N-81	AGATCTTTATTTTAA
723.			
		rb-N-82	TAGATCTTTATTTTTAA
724.		rb-N-83	AATCATCATTAATT
725.		rb-N-84	AAATCATCATTAATT
726.		rb-N-85	AAAATCATCATTAATT
727.		rb-N-86	TAAAATCATCATTAATT
728.	•	rb-N-87	TTAAAATCATCATTAATT
729.		rb-N-88	TTTAAAATCATCATTAATT
730.		rb-N-89	ATTTAAAATCATCATTAATT
731.		rb-N-90	AATTTAAAATCATCATTAA
732.		rb-N-91	
		_	GAATTTAAAATCAT
733.		rb-N-92	TGAATTTAAAATCAT
734.		rb-N-93	TTAAAATAGGAAAT
735.		rb-N-94	AATTTCTCTTTAAA
736.		rb-N-95	AAATTTCTCTTTAAA
737.		rb-N-96	TAAAATTTTGAATG
738.		rb-N-97	CTAAAATTTTGAAT
739.		rb-N-98	TTTGCTAAAATTTT
740.		rb-N-99	ATATGAAAAATGTT
741.		rb-N-100	
			TTTTAAATTAAGCA
742.		rb-N-101	TTGTAAAAATCAAA
743.		rb-N-102	TTTGTAAAAATCAAA
744.		rb-N-103	TTTGATAAAACTTT
745.		rb-N-104	ATGTTTTATCATTT
746.			
		rb-N-105	AATGTTTTATCATTT
747.		rb-N-106	AAATGTTTTATCATTT
748.		rb-N-107	TAAATGTTTTATCATTT
749.		rb-N-108	TCTAAATGTTTTAT
750.		rb-N-109	TTCTAAATGTTTTAT
751.		rb-N-110	TAAGATCAAATAAA
752.		rb-N-111	ATAAGATCAAATAAA
753.		rb-N-112	AATAAGATCAAATAAA
754.		rb-N-113	TAATAAGATCAAATAAA
755.		rb-N-114	
			TTAATAAGATCAAATAAA
<b>756.</b>		rb-N-115	TTTAATAAGATCAAATAAA
757.		rb-N-116	TTGTTTAATAAGAT
758.		rb-N-117	ATTGTTTAATAAGAT
759.		rb-N-118	TGATTGTTTAATAA
760.			
		rb-N-119	TTGATTGTTTAATAA
761.		rb-N-120	TTTGATTGTTTAATAA
762.		rb-N-121	TTTTATAAAACAGT
763.		rb-N-122	TTTTTATAAAACAGT
764.		rb-N-123	TTTTTTATAAAACAGT
765.		rb-N-124	CTTTTTTATAAAACA
766.		rb-N-125	ACTTTTTTATAAAACA
767.		rb-N-126	CACTTTTTTATAAAA
768.		rb-N-127	ACACTTTTTTATAAAA
769.		rb-N-128	
			TACACTTTTTTATAAAA
770.		rb-N-129	ATACACTTTTTTATAAAA
771.		rb-N-130	ATTTTGAATTTAAG
772.		rb-N-131	GATTTTGAATTTAA
773.		rb-N-132	TGATTTTGAATTTAA
774.		· ·	
		rb-N-133	ATGATTTTGAATTTAA
775.		rb-N-134	AATGATTTTGAATTTAA
776.		rb-N-135	ATAATAGAATCATA
. 777.		rb-N-136	TATAATAGAATCATA
778.		rb-N-137	TATAATAGAATCAT
779.		rb-N-138	TACTATAATAGAAT
780.		rb-N-139	ATACTATAATAGAAT
781.		rb-N-140	AATACTATAATAGAAT
782.		rb-N-141	AGAATACTATAATA
783.		rb-N-142	
_			TAGAATACTATAATA
784.		rb-N-143	ATAGAATACTATAATA
Fig.	3 - 12		
9.	) - TZ		

15 / 36 785. rb-N-144 TATAGAATACTATAATA 786. rb-N-145 TTATAGAATACTATAATA 787. rb-N-146 AATATTTGTTTTCA 788. **AAATATTTGTTTTCA** rb-N-147 789. rb-N-148 **AAAATATTTGTTTTCA** 790. rb-N-149 CAAAATATTTGTTTT 791. **AAATTTTATATGGA** rb-N-150 792. rb-N-151 TGAAATTTTATATG 793. rb-N-152 CTGAAATTTTATAT TCTGAAATTTTATAT 794. rb-N-153 795. rb-N-154 TTCTGAAATTTTATAT 796. ATCTGATTTATTTT rb-N-155 797. rb-N-156 **AAGATATTAAATGT** 798. rb-N-157 TGAAGATATTAAAT 799. rb-N-158 ATAAATAACAATGA 800. rb-N-159 TATAAATAACAATGA 801. rb-N-160 GTATAAATAACAAT 802. rb-N-161 TGTATAAATAACAAT 803. rb-N-162 TTGTATAAATAACAAT 804. rb-N-163 TCTTGTATAAATAA 805. rb-N-164 **ATCTTGTATAAATAA** 806. rb-N-165 **AATCTTGTATAAATAA** 807. rb-N-166 ACAACTTTTTAAAT 808. rb-N-167 TACAACTTTTTAAAT 809. rb-N-168 TACAACTTTTTAAA 810. rb-T-1 CGGGGGTTTTGGGCGGCATG 811. rb-T-2 TTTTCGGGGGGTTTTGGGCGGCA 812. rb-T-3 TCGGGGGTTTTGGGCGGC 813. rb-T-4 GGTGGCGGCCGTTTTTCGGGGGGGT 814. CCGGGGGTTCCGCGGCGGCAGCG rb-T-5 815. rb-T-6 CGGGGGTTCCGCGGCGG 816. rb-T-7 GGCGGCGGTGCCGGGGGTTCCGC 817. rb-T-8 GGAGGGGCGCGGCGGTG 818. rb-T-9 GGGGCGCGCGCGCG 819. rb-T-10 GGGGCGGCGCG 820. rb-T-11 AGGGGCCTGGTGGAAG 821. rb-T-12 TAGGGGGCCTGGTG 822. GTAGGGGGCCTGGT rb-T-13 823. rb-T-14 GAGGTATTGGTGACAAGGTAGGGGGC 824. rb-T-15 TCTTCAGGGGTGAAATATAGATGTTC 825. rb-T-16 GGACTCTTCAGGGGTG

Fig. 3 - 13

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826
                   TCGGACTATA CTGC
827
                   CAGTTCGGAC TATACT
828
                   AAGCCTAAGA CGCA
829
                   GCCCAAGTTC AACA
830
                   TGAAAAGTCG CGGT
831
                   GGTTAATTAA GATGCCTC
832
                   TCTCTAAGAG CGCA
833
                   ACGTGAGGTT AGTTTG
834
                   CACGTGAGGT TAGT
835
                   CATAGAACAG TCCG
                   CAGTCATAGA ACAGTC
836
837
                   CTTTGCAGTC ATAGAACA
838
                   TGCAGTCATA GAAC
839
                   GGTCGTTTCC ATCT
840
                   CATAGAAGGT CGTTTC
841
                   CGTCATAGAA GGTC
842
                   CATCGTCATA GAAGG
843
                   GGACGGGAGG AACGAGGCGT TGAG
844
                   TAGCCATAAG GTCC
845
                   GGTTACTGTA GCCA
846
                   GGTTACTGTA GCCA
847
                   AGTTCTTGGC GCGGAGGT
848
                   AGGTGAGGAG GTCCGAGT
849
                   TGGACTGGAT TATCAG
850
                   GTGGTGGTGA TGTGCCCG
851
                   TGTCACGTTC TTGG
852
                   CTCATCTGTC ACGT
853
                   CGAAGCCCTC GGCGAACC
854
                   GCGTGTTCTG GCTGTGCAGT TCGG
855
                   CTGCCCCGTT GACC
856
                   AGGTTTGCGT AGAC
857
                   GGTTGAAGTT GCTG
858
                   CTGGGTTGAA GTTG
859
                   TGCTGCACGG GCATCTGCTG
860
                   GGCACTGTCT GAGGCTCCTC CTTCAGG
861
                   ACTCCATGTC GATG
862
                   CTCTCCGCCT TGATCC
863
                   GTTCCTCATG CGCTTC
864
                   CTGAGCTTTC AAGG
865
                   GCGATTCTCT CCAGCTTCCT TTTTCG
866
                   CTGAGCTTTC AAGGTTTTCA CTTTTTCCTC
867
                   TCCCTGAGCA TGTT
868
                   TCTGTTTAAG CTGTGC
869
                   CTTTCTGTTT AAGCTGTG
870
                   GGTTCATGAC TTTCTG
871
                   CGTGGTTCAT GACT
872
                   ACTGTTAACG TGGTTC
873
                   CCACTGTTAA CGTG
874
                   CCCACTGTTA ACGT
875
                   AGCATGAGTT GGCA
876
                   GCGTTAGCAT GAGT
877
                   GTTTGCAACT GCTG
878
                   CAAAATGTTT GCAACTGC
Fig.
      4 - 1
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879
                   TCCATTTTAG TGCACATC
880
                   CTGTTCCATT TTAGTGCA
881
                   GTGTATGAGT CGTC
                   CTGTGTATGA GTCG
882
883
                   CGTAGCTGTG TATG
884
                   TCGTGTAGAG AGAG
885
                   AGTTTGTAGT CGTGTAGA
886
                   GTTTGTAGTC GTGTAG
887
                   AGTTTGTAGT CGTG
888
                   GGAGTTTGTA GTCG
889
                   TCAGGAGTTT GTAGTC
890
                   GTTTCAGGAG TTTGTAGT
891
                   TCGGTTTCAG GAGT
892
                   TTGAGACTCC GGTA
893
                   ACCAGAAAAG TAGCTG
894
                   CCTGACCAGA AAAG
895
                   ATTCAGGCGT TCCA
896
                   GGTAAAAGTA CTGTCC
897
                    GGGTAAAAGT ACTGTC
898
                    GCACCTCCAC CGCTGCCA
899
                    CTCCTGCTCC TCGGTGAC
900
                    GCTTTGACAA AGCC
901
                    CTTGTGCAGA TCGT
902
                    TCATCTTGTG CAGATC
903
                    GTTCATCTTG TGCAGA
                    CGTGGTTCAT CTTG
904
905
                    TCACGTGGTT CATC
906
                    GGTTGGTGTA AACG
907
                    TACGAGCTCC CGGTCCCGAC
908
                    TAGCTGATGG TGGT
909
                    TCCTTGAAGG TGGA
910
                    TCTTCCATGT TGATGG
911
                    CTTTGATGCG CTCT
912
                    CTCCACTTTG ATGC
913
                    GCTCCAGCTT CCGCTTCCGG CACTTGGTGG
914
                    GGCCTTGAGC GTCTTCACCT TGTCCTCCAG
915
                    TGACCTTCTG TTTGAG
916
                    CATGACCTTC TGTTTG
917
                    GTCATGACCT TCTG
918
                    CGAGAACATC ATCG
919
                    GTAGTCTGCG TTGA
920
                    GCTGCAGCGG GAGGATGACG
921
                    AGTAAGAGAG GCTATC
922
                    GTAGTAAGAG AGGC
923
                    GGTAGTAAGA GAGG
924
                    GTGAGTGGTA GTAAGA
925
                    GTCCGTGCAG AAGTCCTG
926
                    GAATGAAGTT GGCACT
927
                    GGAATGAAGT TGGC
928
                    GGGAATGAAG TTGG
929
                    GCTGCACCAG CCACTGCAGG TCCGGACTGG
930
                    TCATGGTCTT CACAAC
931
                    CAATGCTCTG CGCTCGGCCT CCTGTCATGG
Fig.
```

	932	CTAGAGTTCC	TICA C	
	933	GAGTACGCTA		
	934	GAAGAGTACG		
	935		CCCAGCCCCC	A CA MMCCC
	936		GTACTGGGCT	ACATICCC
	937			
	938	GTTACGGATG		
	939	CAGTTACGGA		
	940	CCAGTTACGG		
	941	AGAGTCTGAG		
		GTGAGACTCA		
	942	TCTTAGGGTG		
	943	GAGAGTACTT		
	944	GGAAGAAACT	-	
	945	CTTAGGGAAG		
	946	CGGTAAGAAA		
	947	AGCATGCGGT		
	948	GTCTGAAAGC		
	949	AGAACAAAGA	AGAGCC	
	950	CAAGAGAACA		
	951	CAGCAAGAGA	ACAAAG	
	952	TCCTCAGCAA		
	953	AGGTGTGACT		
	954	GAATAGGTGT		
	955	CAGAATAGGT		
	956	GCAGAATAGG		
	957	CAGTTGCAGA		
	958	GAAACCATTT		
	959	TGTGAAACCA		
	960	CACTGTGAAA		
	961	CCACTGTGAA		
	962		CCTGCAGCTT	CCCTGCTTCC
	963	CACCTCCATT		
	964	CAGTAAAAGT		
	965	CGACATTCAG		
	966 967	GACCGACATT		
	968	CTTCTGGAGA		
	969	CATCTTATTC		
		CAGCCATCTT		
	970	TGCAGCCATC		
	971 972	GAGTGTATCA		
		GGAGTGTATC	-	
	973	CTTGGAGTGT		
	974 975	ACAGAGTACC		
	976	CCAACTTTCC		
	977	CCTTATGCTC		
	977	GTCTTACTCA		
	979	ACAGTCTTAC		
	980	CATAAGACAC		
	981	GAAAGCATAA	· —	
	982	GGAAAGCATA		
	983	AGGGATAAAG CCTGTATACA		
	984	TGTCTCCTGT		
•		19101001	DAJMIN	
	Fig. 4 - 3			
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985	CATCTTCTAG TTGGTC
986	CTCATCTTCT AGTTGG
987	CTTCTCATCT TCTAGTTG
988	CAAAGCAGAC TTCTCA
989	CTGCAAAGCA GACT
990	CTAGTTTTTC CTTCTCCT
991	TCTAGTTTTT CCTTCTCC
992	CAGGATGAAC TCTAGT
993	TCGTAGAAGG TCGT
994	AGGGTTACTG TAGC
995	GTAGTGGTGA TGTG
996	CGTCGTAGAA GGTC
997	TTTCGTGCAC ATCC
998	AGTTTGTAGT CGTGAAGA
999	CGAGAACATC ATGG
1000	GTAGTAGGAA AGGC
1001	GGTAGTAGGA AAGG
1002	GGAATGGTAG TAGG
1003	GGTCATTGAG AAGAG
1004	GCTAATGTTC TTGACC
1005	GCCAAGGTCCTCAT
1005	GGAGTCTATCTCCA
1007	CCAAAGAATCCTGACT
1007	CACATGCTTAGTGG
1009	CTCGTAAATGACCG
1010	AGGAATCTCGTAAATGAC
1011	CAGCAGCGATTCAT
1012	GGAGATCATCAAAGGA
1013	CTCAGCAATGGTCA
1014	GATCTCGAACACCT
1015	CACAATCTCGATCTTTCT
1016	CCTTCTTAAAGATTGGCT
1017	CACATACCAACTGG
1018	AGCTTGATGTGAGG
1019	GAAGTTGTAGCTTGATGT
1020	GCTTGAAGTTGTAGCT
1021	CTGCTTGAAGTTGTAG
1022	GACACAACTCCTCT
1023	TCCTTTGATAGACACAAC
1024	CTCGTTTGATAGACAC
1025	GGTTAGCACACT
1026	GGTAACGGTTAGCA
1027	CGTAACACATTTAGAAGC
1028	CTCATCCGTAACAC
1029	CCGGTAAGTATTGTAGTT
<b>10</b> 30	GGTGTATTTCCTTGAC
1031	ACATACCAACTGGTGT
1032	GTCCCTATACGAAC
1033	TTCATGTCTG TGCC
1034	GTAGGTGAGT TCCA
1035	GTTGTGAGCG ATGA
1036	CATAGTTGTC CTCAAAGA
1037	GGCATAGTTG TCCT
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Fia.	4 - 4

Fig. 4-4

**WO 98/33904** 

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1038	CATTGTCTAG	CACG
1039	CTCCATTGTC	
1040	GTATTGTTCA	
1041	TCAAGATCTC	
1042	CACAAAATCG	
1043	TCCTTCCACA	
1044	GTGGAAGATG	
1045	TCTTGTGGAA	
1046	TCTATCAGTG	
1047	GGTTGGTGTC	
1048	ACATCGGAGA	
1049	CCTTACACAT	
1050	ACAATCCTCA	
1050		
1051	GCTCTGACAA	
	TGGTTGAAGT	
1053	CTGTGGTTGA	
1054	GTTGTAGGTG	ACCA
1055	CTGTGTTGTA	<del></del>
1056	GACTCAAACG	TGTC
1057	CATGGACTCA	
1058	CGAATGTATA	
1059	CCGAATGTAT	
1060	GCCGAATGTA	-
1061	GTAGTTGTAG	GGAC
1062	TAGAAAGGTA	GTTGTAGG
1063	GTAGAAAGGT	AGTTGTAG
1064	CGTAGAAAGG	TAGTTG
1065	CCGTAGAAAG	GTAG
1066	GACCATAGCA	CACT
1067	GGATATTGGC	ACTG
1068	CCTGGATATT	GGCA
1069	GCTCCCAAAG	ATCT
1070	CCCATCAAAG	CTCT
1071	CAAACACTTG	GAGC
1072	GTCTCAAACA	CTTGGA
1073	GAGTCTCAAA	CACTTG
1074	GTAACCTGTG	ATCTCT
1075	GGTAACCTGT	GATC
1076	GTATAGGTAA	CCTGTG
1077	TGAGATGTAT	AGGTAACC
1078	TGCTGAGATG	TATAGG
1079	CCATGCTGAG	ATGT
1080	GGATTACTTG	CAGG
1081	TGTTATGGTG	GATGAG
1082	GGTGTTATGG	TGGA
1083	GCAGTTGACA	CACT
1084	AGTACTCGGC	ATTC
1085	CATTCACATA	CTCCCT
1086	TCCAAAACAG	
1087	GGTCCTTATA	
1088	CAGAATGCCA	
1089	ACGAGAATGC	
1090	GATCCCAAAG	
		~ * <del>~</del> ~ ~ *
Fig. 4 - 5		

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1091		TCGCTTGATG	AGGA
1092		CATCGTGTAC	TTCC
1093		GCATCGTGTA	CTTC
1094		ACTGTGCCAA	AAGC
1095		CTTGTAGACT	GTGC
1096		CCCTTGTAGA	CTGT
1097		TCAACACTTT	GATGGC
1098		CCCTCAACAC	TTTG
1099		GTGTTTTCCC	TCAACA
1100		GTATGCTTCG	TCTAAG
1101		CGTATGCTTC	GTCT
1102		CCATCACGTA	TGCT
1103		GCATAAGCTG	TGTC
1104		CATGGTCTAA	GAGG
1105		CAATCTGCAT	ACACCA
1106		GGCAATCTGC	ATAC
1107		CTGTCTCGTC	AATG
1108		CATAACTCCA	CACATC
1109		AGTCACACCA	TAACTC
1110	•	ACAGTCACAC	CATAAC
1111		CCCCAAAAGT	CATC
1112		TCGTAAGGTT	TGGC
1113		GATCCCATCG	TAAG
1114		CAATGGTGCA	GATG
1115		GACATCAATG	GTGC
1116		GTAGACATCA	ATGGTG
1117		CATGATCATG	TAGACATC
1118		CCATGATCAT	GTAGAC
1119		CATTTGACCA	TGATCATG
1120		CCAACATTTG	ACCATG
1121		TCATCCAACA	TTTGACCA
1122		GAGTCAATCA	TCCAACAT
1123		CAGAGTCAAT	CATCCA
1124		CCGACATTCA	GAGT
1125		GAATTCAGAC	ACCAAC
1126		GATGACCACA	AAGC
1127		CCATCAAATA	CATCGG
1128		TCACCATCAA	ATACATCG
1129		CAACGTAGCC	ATCA
1130		ACGTCTTTGA	CGAC
1131		CAAAAACGTC	TTTGACGA
1132		GGCAAAAACG	TCTTTG
1133		CAAAGGCAAA	AACGTC
1134		GTGTCAAGTA	CTCG
1135		GTAATAGAGG	TTGTCG
1136		CCCAGTAATA	GAGG
1137		CATGGTGCTC	ACTG
1138		GTGCCTGTAC	GTAC
1139		TGCAGGTGGA	TAGT
1140		CATGTCGATA	GTCTTGCA
1141		GTCGATAGTC	TTGC
1142		CCATGTCGAT	AGTC
1143		CTCCATGTCG	ATAG
Fig.	4 - 6		
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		•
1144	CTTGGACAGG	ATCT
1145	TGCTGTTGTA	CAGG
1146	GTGCTGTTGT	ACAG
1147	TTGGCGTAGT	AGTC
1148	TCCACCATTA	GCAC
1149	GATTTCGTTG	TGGG
1150	GTCATAGATT	TCGTTGTG
1151	TGTACTCTGC	TTGAAC
1152	GTGTACTCTG	CTTG
1153	TGCTGTGTGT	ACTC
1154	CTGATGTGTT	GAAGAACA
1155	CTCTGATGTG	TTGAAG
1156	GCTCTGATGT	GTTG
1157	GAGCTCTGAT	GTGT
1158	CACTTTTAAC	TTGAGCCT
1159	CTCCACTTTT	AACTTGAG
1160	TGCTGTATTT	CTGGTACA
1161	CCAGGAATTG	
1162	TTGCTGAGGT	ATCG
1163	GATAACCACT	
1164	CAAAAGATAA	
1165	CGGTGACATC	
1166	CCTCAATTTC	<del>_</del>
1167	GTTATCCCTG	
1168	GCAGTGTGTT	
1169	GATGTCCACT	
1170	TAGTGAACCC	
1171	TGCCATGAAT	
1172	GTTCATGCCA	
1173	CATGAGAAGC	<del>_</del>
1174	GCTTTGCAGA	
1175	GAGCTTTGCA	
1176	TAGTTGGTGT	
1177	CTGAAGCAAT	
1178	AGCTGAAGCA	
1179	GGAGCTGAAG	
1180	CAATGTACAG	
1181	GGAAGTCAAT	
1182	CGGAAGTCAA	
1183	GCGGAAGTCA	V j
1184	AGTTGGCATG	
1185	GCAGAAGTTG	
1186	CTCCAAATGT	
1187	ACCTTGCTGT	
1188	TGCTGGTTGT	<u>-</u>
1189	GGTTATGCTG	
1190	GTAGTACACG	
1191	CGTAGTACAC	
1192	CACGTAGTAC	
1193	CATGTTGGAC	
1194	GCACGATCAT	
1195	CACACAGTAG	
1196	GATCAGAAAA	
Fig. 4 - 7		

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1197	ACCGTGACCA GATG
1198	GTAGACAGGC TGAG
1199	TATCGAGTGT GCTG
1200	TTGCGCATGA ACTG
1201	TTGCTCAGGA TCTG
1202	ACTGGTGAGC TTCA
1203	GCTCAGGATA GTCT
1204	TGTAGATGGA AATCACCT
1205	TGGTGCTGTT GTAG
1206	TTCTCCTGGA GCAA
1207	TACTCTTCGT CGCT
1208	CTTGGCGTAG TACT
1209	CGGCATGTCT ATTTTGTA
1210	CGGGATGGCA TTTT
1211	CTGTAGAAAG TGGG
1212	ACAATTCTGA AGTAGGGT
1213	ATTGCTGAGA CGTCAAAT
1214	TCTCCATTGC TGAG
1215	TCACCAAATT GGAAGCAT
1216	CTCTGAACTC TGCT
1217	AACGAAAGAC TCTGAACT
1218	TGGGTTCTGC AAAC
1219	CTGGCTTTTG GGTT
1220	GTTGTTCAGG CACT
1221	TCTGATATAG CTCAATCC
1222	TCTTTGGACT TGAGAATC
1223	TGGGTTGGAG ATGT
1224	TGCTGTCGAT GTAG
1225	ACAACTTTGC TGTCGA
1226	ATTCGCCTTC TGCT
1227	GAAGGAGAC CATT
1228	TCAGTTACAT CGAAGG
1229	TGAAGCCATT CATGAACA
1230	TCCTGTCTTT ATGGTG
1231	AAATCCCAGG TTCC
1231	GGACAGTGTA AGCTTATT
1232	GTACAAAAGT GCAGCA
1233	TAGATGGTAC AAAAGTGC
1234	CACTTTTATT TGGGATGATG
1235	GCAAATCTTG CTTCTAGT
1230	GTGCCATCAA TACC
1237	GGTATATGTG GAGG
	TCTGATCACC ACTG
1239	TCTGATCACC ACTG  TCCTAGTGGA CTTTATAG
1240	TTTTTCCTAG TGGACT
1241	CAATAACATT AGCAGG
1242	
1243	AAGTCTGTAG GAGG
1244	TCTGTTGTGA CTCAAG
1245	GTTGGTCTGT TGTG
1246	CAAAGCACGC TTCT
1247	TTTCTAAAGC AATAGGCC
1248	GCAATTATCC TGCACA
1249	ACGTAGGCAG CAAT
Fig.	4 - 8

1250	ATCAATGTAA	AGTGGACG
1251	CTAGATCCCT	CTTG
1252	CCATTTCCAC	CCTA
1253	TGGGTTCGTG	TATC
1254	TGGCATTGTA	CCCT
1255	TCCAGCACAG	AAGT
1256	ATAAATACGG	GCATGC
1257	AGTGTCTGAA	CTCC
1258	TGTGCTGAGT	GTCT
1259	ATAAGCTCAG	GACC
1260	AGGAGAAGCA	GATG
1261	AGCAAGGAGA	AGCA
1262	AATCTTGGGA	CACG
1263	TAGAGAATGG	TTAGAGGT
1264	GTTTTGCCAA	TGTAGTAG
1265	CTTGGGTGTT	TTGC
1266	GCAAGACTTT	ACAATC
1267	GCATTTGCAA	GACTTTAC
1268	TTTAGCTGCA	TTTGCAAG
1269	GCCACTTTTC	CAAG
1270	TTGGTCTTGC	CACT
1271	CAGCACACAG	TAGT
1272	CGATAGTCTT	GCAG

	mar 00 14/1	25	CTTTCACCAAATTGGAAG
1273	TGF-\(\beta^2 - 14/1\)		
1274	TGF-\(\beta^2 - 14/2\)		CACCAAATTGGAAGC
1275	TGF-ß2-14/3		TCACCAAATTGGAAGC
1276	TGF-ß2-15/1		CTCTGGCTTTTGGG
1277	TGF-ß2-9/1		CGGCATGTCTATTTG
1278	relA-1		CACTACAGACGAGC
1279	relA-2		CGTGCACTACAGACG
1280	relA-3		GGAACAGTTCGTCC
1281	relA-4		GAACAGTTCGTCCATG
1282	relA-5		CCAGAGTTTCGGTTC
1283	relA-6		CTAGGACTGGGACAG
1284	relA-7		CGCACTTGTAGCG
1285	relA-8		CTCGCACTTGTAGC
1286	relA-9		GCACTTGTAGC
1287	relA-10		GCGCACTGTCCCTG
1288	relA-11		CCAGGGAGATGCGC
1289	relA-12		GCCGGTGAGGAGG
1290	relA-13		CCGGTGAGGAGGG
1291	relA-14		CGGTTCACTCGGC
1292	relA-15		GAGTTTCGGTTCACTC
1293	relA-16		GGCACGATTGTCAAAG
1294	relA-17		CAGGCGTCACCCCC
1295	relA-18		GCAGGCGTCACCC
1296	p105/p50-1		CTCCCTCCTAAGC
1297	p105/p50-2		CCCTCCTAAGCGG
1298	p105/p50-3		CGAGTCCGCGTTCG
1299	p105/p50-4		CATCTTCTGCCATTC
1300	p105/p50-5		GTGTTTTCCCACCAG
1301	p105/p50-6		GGTTTTGGTTCACTAG
1302	p105/p50-7		GCATCTTCACGTCTCC
1303	p105/p50-8		CTTCACGTCTCCTGTC
1304	p105/p50-9		GTCACCGCGTAGTC
1305	p105/p50-10		CAAATAGGCAAGGTC
1306	p105/p50-11		CTTGCAAATAGGCAAG
1307	p105/p50-12		TGCTTGCAAATAGG
1308	p105/p50-13		CTGCTTGCAAATAGG
1309	p105/p50-14		GCAGGTGGATATTT
1310	p105/p50-15		CTGCTGTTGGCAG
1311	p105/p50-16		CACTAGTTTCCAAGT
1312	p105/p50-17		GTTTTGGTTCACTAG
1313	p105/p50-18		CTTTGATTTCAGGATAG
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Fig. 5 - 1

		26 / 36
1314	p105/p50-19	GCACTTCTTCTTATCT
1315	p105/p50-20	CCAAGTCAGATTTCC
1316	p105/p50-21	GTTTCCAAGTCAGATTTC
1317	p105/p50-22	GGTTCACTAGTTTCC
1318	p105/p50-23	GGTTTTGGTTCACTAG
1319	p105/p50-24	CCGAAAAATTGGGCA
1320	p105/p50-25	CCGAAAAATTGGG
1321	p105/p50-26	CTATCCGAAAAATTGG
1322	p105/p50-27	GTTGATAATGTCATCAG
1323	p105/p50-28	CTCATGTTGATAATGTC
1324	p105/p50-29	CIGTCACCGCGTAG
1325	p105/p50-30	CGTCTCCTGTCACCG
1326	p105/p50-31	CTTCACGTCTCCTG
1327	p105/p50-32	GAGAACTTATCATGTC
1328	p105/p50-33	GCTATATGCAGGG
1329	p105/p50-34	CCAGCTGCTATATGCAGG
1330	p105/p50-35	AGGCTAAATTTTGCCT
1331	p105/p50-36	GGCTAAATTTTGCC
1332	p105/p50-37	GGCTAAATTTTGCCTTC
1333	p105/p50-38	GCAGGCTAAATTTTGCC
1334	p105/p50-39	GAGTTACCCAAGCG
1335	p105/p50-40	CAGAGTTACCCAAGCG
1336	p105/p50-41	CAGAGTTACCCAAG
1337	p105/p50-42	ACAGAGTTACCCAAG
1338	p105/p50-43	GGTGCAAAACAGAG
1339	p105/p50-44	CTAGGTGCAAAACAG
1340	p105/p50-45	GAGAACTTTATCATGTCC
1341	p105/p50-46	GCTAGATGAATGGC
1342	p105/p50-47	GCAAACATGGCAGGC
1343	p105/p50-48	CAGCAAACATGGCA
1344	p105/p50-49	GCAGCAAACATGGC
1345	p105/p50-50	AGCAGCAAACATGG
1346	p105/p50-51	CAGCAGCAAACATG
1347	p105/p50-52	AGCAGCAGCAAACA
1348	p105/p50-53	CAGCAGCAGCAAACA
1349	p105/p50-54	CAGCAGCAGCAAAC
1350	p105/p50-55	CACCAGCAGCA
1351	p105/p50-56	GCATTGACGTCAGC
1352	p105/p50-57	GATGTTGTCGTGCTC
1353	p105/p50-58	TGAGATGTTGTCGTGCT
13 <i>5</i> 4	p105/p50-59	TGAGATGTTGTCGTG

Fig. 5 - 2

		27 / 36
1355	p105/p50-60	GCCAATGAGATGTTG
13 <i>5</i> 6	p105/p50-61	CTGCCAATGAGATG
13 <i>5</i> 7	p105/p50-62	CACATGGGCATCAC
1358	p105/p50-63	TGTCCACATGGGCA
1359	p105/p50-64	GTACTGTCCACATG
1360	p105/p50-65	CAGCTGCTATATGC
1361	p105/p50-66	GTTCTCCACCAGGG
1362	p105/p50-67	AGTTCTCCACCAGG
1363	p105/p50-68	CAAAGTTCTCCACCAG
1364	p105/p50-69	CCAAGAGTCATCCAGG
1365	p105/p50-70	CCCAAGAGTCATCC
1366	p105/p50-71	CCTGCATTTTCCCAAG
1367	p105/p50-72	TCCTGCATTTTCCC
1368	p105/p50-73	GCCATATCTAGAGGC
1369	p105/p50-74	TCACATCTTCAGCC
1370	p105/p50-75	GCTTCACATCTTCAGC
1371	p105/p50-76	CAGCTTCACATCTTC
1372	p105/p50-77	GTAACTTATACAGCTGC
1373	p105/p50-78	CCAGTTTTTGTCTGG
1374	p105/p50-79	CCATTTGTCTCAGG
1375	p105/p50-80	GTGTAGCCCATTTG
1376	p105/p50-81	GCTTCGGTGTAGCC
1377	p105/p50-82	GATCACITCAATTGCTTC
1378	p105/p50-83	CTTGTGGAGGCAGG
1379	p105/p50-84	GCTGCCTTGTGGAG
1380	p105/p50-85	CTATTTGCTGCCTTGTGG
1381	p105/p50-86	GGATGTCTCCACGC
1382	p105/p50-87	GGAAGGATGTCTCC
1383	p105/p50-88	TGCGGAAGGATGTC
1384	p105/p50-89	GTTTGCGGAAGGATGTC
1385	p105/p50-90	GCTGAGTTTGCGGA
1386	p105/p50-91	GGTAAAGCTGAGTTTG
1387	p105/p50-92	TCGGTAAAGCTGAG
1388	p105/p50-93	GACTCGGTAAAGCTG
1389	p105/p50-94	AGAGACTCGGTAAAGC
1390	p105/p50-95	GAAATTGTCAGCAGGC
1391	p105/p50-96	GAAATTGTCAGCAGG
1392	p105/p50-97	GGAAATTGTCAGCAGG
1393	p105/p50-98	GGAAATTGTCAGCAG
1394	p105/p50-99	GGGAAATTGTCAGC
1395	p105/p50-100	GTGTGGGAAATTGTC
	L L	

		28 /	36	
1396	p105/p50-101			GGTTTACACGGTGTG
1397	p105/p50-102			GCTTTGGTTTACACG
1398	p105/p50-103			GCACCTTTGGGATGC
1399	NFKB2-1			CCAGGTTCTGCTTCC
1400	NFKB2-2			GCTCTGTCTAGTGGC
1401	NFKB2-3			ACTCTCCATGTCTC
1402	NFKB2-4			CAACTCTCCATGTCTC
1403	NFKB2-5			CAACTCTCCATGTC
1404	NFKB2-6			AGCAACTCTCCATG
1405	NFKB2-7			GTAGCAACTCTCCATG
1406	NFKB2-8			GTAGCAACTCTCCA
1407	NFKB2-9			GGTTGTAGCAACTCTCC
1408	NFKB2-10			CGGGCAGTCCTCCA
1409	NFKB2-11			GCACCGGGCAGTC
1410	NFKB2-12			AGGCACCGGGCAG
1411	NFKB2-13			GTGTGTTACCAGGTC
1412	NFKB2-14			TGTGTGTTACCAGGT
1413	NFKB2-15			TGGGTCACTGTGTG
1414	NFKB2-16			CAGACTGTGGGCATG
1415	<b>NFKB2-17</b>			CCCACCAGACTGTGGG
1416	NFKB2-18			CCACCAGACTGTGG
1417	NFKB2-19			TGCCCACCAGACTG
1418	NFKB2-20			CGGCTTCCTCCC
1419	NFKB2-21			CCTTGTCTTCCACC
1420	<b>NFKB2-22</b>			ACCGAGGCTGCCAC
1421	NFKB2-23			GGAAGAAACCGAGG
1422	<b>NFKB2-24</b>			GGGAAGAAACCGAG
1423	NFKB2-25			GGCCATCTGCGCC
1424	NFKB2-26			GCGGCCATCTGCG
1425	<b>NFKB2-27</b>			GTGGCGGCCATCTG
1426	NFKB2-28			ACCGTGGCGGCCAT
1427	NFKB2-29			GCCGCTCAATCTTCATC
1428	NFKB2-30			CTTCATCTTGTGATAGG
1429	NFKB2-31			GCTCAATCTTCATCTTG
1430	NFKB2-32			CAGAAACACTGTTACAG
1431	<b>NFKB2-33</b>			CAGTTGCAGAAACACTG
1432	NFKB2-34			GTTTCAGTTGCAGAAAC
1433	NFKB2-35			CTTCCACCAGAGGG
1434	NFKB2-36			GTCTTCCACCAGAG
1435	NFKB2-37			CTTGTCTTCCACCAGAG
1436	NFKB2-38			TCCTTGTCTTCCAC

Fig. 5 - 4

		29 / 36
1437	NFKB2-39	CTTCCTTGTCTTCCAC
1438	NFKB2-40	CATCTTGTGATAGGG
1439	NFKB2-41	GCTAGGTGCAGTGGT
1440	NFKB2-42	GATGGCTAGGTGCA
1441	NFKB2-43	GTGGATGATGCTAG
1442	NFKB2-44	CCCGTGGATGATGG
1443	NFKB2-45	CTGCCCGTGGATGA
1444	NFKB2-46	AGAGCCTCCACCCA
1445	NFKB2-47	GTTGTACTCTCGAGC
1446	NFKB2-48	CGTTGTACTCTCG
1447	NFKB2-49	CGCGTTGTACTCTC
1448	NFKB2-50	GAGTCTCCATGCCG
1449	NFKB2-51	CTGAGTCTCCATGC
1450	NFKB2-52	CATGGCTGAGTCTC
1451	NFKB2-53	TGCATGGCTGAGTC
1452	NFKB2-54	GCGTTCACGTTGGC
1453	NFKB2-55	GTGCGAGCGTTCAC
1454	NFKB2-56	AGGTGCGAGCGTTC
1455	NFKB2-57	GCAAAGGTGCGAGC
1456	NFKB2-58	CCTGGTGGCTCAGG
1457	NFKB2-59	GTCAGTCACCTGAG
1458	NFKB2-60	CAGGTCAGTCACCTG
1459	NFKB2-61	CAGCAGGTCAGTCAC
1460	NFKB2-62	GCAGCAGGTCAGTC
1461	NFKB2-63	CATTTAGCAGCAAGGTC
1462	NFKB2-64	GCAGCATTTAGCAGC '
1463	NFKB2-65	CTGAGCAGCATTTAG
1464	NFKB2-66	CCCATGAGAATCCT
1465	NFKB2-67	CCTTCCCATGAGAATCC
1466	NFKB2-68	TCCTCCCCTTCCCA
1467	NFKB2-69	GCCTCCAGTAGACC
1468	NFKB2-70	GTCAGACAGGGCCT
1469	NFKB2-71	CCATGTCAGACAGG
1470	NFKB2-72	GGCCCATGTCAGAC
1471	TANK-1	GCTATTCCTGAAATCAC
1472	TANK-2	CCTCTTGTCTTACC
1473	TANK-3	GGAGAAGAAACCTCTTG
1474	TANK-4	CCTTGCTGAAGTTTCTT
1475	TANK-5	CCAAGACTCCTTGC
1476	TANK-6	CCCTTTCATGGAGC
1477	TANK-7	CCTCTTGGTGTGAC

Fig. 5 - 5

	30 / 36	
1478	TANK-8	GACTAAGGATGCCG
1479	TANK-9	GTGGCAGGACTAAGG
1480	TANK-10	AGACGTGGCAGGAC
1481	I-kappa-Bepsilon-1	CTTCCAGCAGGCAG
1482	I-kappa-Bepsilon-2	GTTCCTCTGCCTGG
1483	I-kappa-Bepsilon-3	GATGTTCCTCTGCCTG
1484	I-kappa-Bepsilon-4	GAGATGTTCCTCTGCC
1485	I-kappa-Bepsilon-5	GTGAGATGTTCCTCTG
1486	I-kappa-Bepsilon-6	CAGAGAGTGAGATGTTCC
1487	I-kappa-Bepsilon-7	CCAGAGAGTGAGATGTTC
1488	I-kappa-Bepsilon-8	GGTCCAGAGAGTGAG
1489	I-kappa-Bepsilon-9	GAGGTCCAGAGAGTG
1490	I-kappa-Bepsilon-10	GGTCCTGTAGTGCC
1491	TRAF-6-1	GATTTTATGATGCAGGC
1492	TRAF-6-2	GACCTGCATCCCTTATTG
1493	TRAF-6-3	TAGTTGATTTTCCAGCAG
1494	TRAF-6-4	GAATCTCACGTTTTGC
1495	TRAF-6-5	CAGAGAAAGAATCTCACG
1496	TRAF-6-6	TTTCACCATCAGAGAAAG
1497	TRAF-6-7	CATTTGGACATTTCACC
1498	TRAF-6-8	CCTTCATTTGGACATTTC
1499	TRAF-6-9	CAATGTGCTTGATGATCC
1500	Rank-1	CGCATCGGATTTCTC
1501	Rank-2	CAAACCGCATCGGATTTC
1502	Rank-3	GAACTGCAAACCGC
1503	Rank-4	GCAGAGAAGAACTGC
1504	Rank-5	GCAAGTAAACATGGG
1505	Rank-6	GGTCCACGTTTTGG
1506	Rank-7	GCAAGGGTCCACGTTT
1507	Rank-8	TGGCTTCTTCTTCAGGG
1508	Rank-9	TCCTGCTGGCTTCTTC
1509	Rank-10	GTCCTGCTGGCTTC
1510	IL-5-1	GGTAGTCTAGGAATTGG
1511	IL-5-2	CTTGCAGGTAGTCTAGG
1512	IL-5-3	GAAACTCTTGCAGGTAG
1513	IL-5-4	CACCAAGAAACTCTTGC
1514	IL-5-5	CATTACACCAAGAAACTC
1515	IL-5-6	CTCGGTGTTCATTACACC
1516	IL-5-7	CTTTCTATTATCCACTCG
1517	IL-5-8	CCAGTTTAGTCTCAACTT
1518	IL-5-9	AACCAGTTTAGTCTCAAC
<b>.</b>		

1519	IL-5-10	ACAAACCAGTTTAGTCTC
1520	IL-13-1	CTCGCGAAAAAGTTTCTT
1521	IL-13-2	CCCTCGCGAAAAAGTTTC
1522	IL-13-3	GTCCCTCGCGAAAAAG
1523	IL-13-4	CAGTTGAACCGTCCC
1524	IL-13-5	GCTTTCGAAGTTTCAGTT
1525	IL-13-6	GATGCTTTCGAAGTTTC
1526	IL-13-7	CTGTCTCTGCAAATAATG
1527	IL-15-1	CACTTATTACATTCACCC
1528	IL-15-2	TTTTCCTCCAGTTCCTC
1529	IL-15-3	GGACAATATGTACAAAACTC
1530	IL-15-4	GTTGATGAACATTTGGAC
1531	IL-15-5	GTGTTGATGAACATTTGG
1532	I-kappaB(newmember)-1	CAAAATTTGGCCAGGG
1533	I-kappaB(newmember)-2	GCCCAAAATTTGGCC
1534	I-kappaB(newmember)-3	CCCAGCCCAAAATTTGG
1535	I-kappaB(newmember)-4	GTCCCCAGCCCAAAATT
1536	I-kappaB(newmember)-5	AAATCGCCAGAGGCTG
1537	I-kappaB(newmember)-6	ACCAAATCGCCAGAGG
1538	I-kappaB(newmember)-7	CATCACCAAATCGCCAG
1539	Prostaglan.Rec.EP3-1	TAGGAGTGGTTGAGGC
1540	Prostaglan.Rec.EP3-2	GTGTAGGAGTGGTTGAG
1541	Prostaglan.Rec.EP3-3	CTGTGTAGGAGTGG
1542	Prostaglan.Rec.EP3-4	CCCACATGCCTGTG
1543	Prostaglan.Rec.EP3-5	CGATGAACAACGAG
1544	Prostagian.Rec.EP3-6	CTGGCGATGAACAACG
1545	Prostaglan.Rec.EP3-7	CGCTGGCGATGAAC
1546	Prostaglan.Rec.EP3-8	GAGCTAGTCCCGTTG
1547	Prostagian.Rec.EP3-9	GCGAAGAGCTAGTCC
1548	Prostaglan.Rec.EP3-10	CCAGTTATGCGAAGAGC
1549	Prostaglan.Rec.EP3-11	CCCCAGTTATGCGAAG
1550	PresenilinI-1	CACATGCTTGGCGC
1551	PresenilinI-2	GATCACATGCTTGGCG
1552	PresenilinI-3	GACAAAGAGCATGATCAC
1553	PresenilinI-4	GAGTCACAGGGACAAAG
1554	PresenilinI-5	GAGAGTCACAGGGAC
1555	PresenilinI-6	GCAGAGAGTCACAGG
1556	PresenilinI-7	CCATGCAGAGAGTC
1557	PresenilinI-8	CCACCATGCAGAGAG
1558	PresenilinI-9	TAGCCACGACCACC
1559	PresenilinI-10	GATTAGCTGCCCATCCTT

1560	PresenilinI-11	GGTATAGATTAGCTGCC
1561	PresenilinI-12	GTATCTTCTGTGAATGGG
1562	PresenilinI-13	CTGGCCCACAGTCT
1563	PresenilinI-14	CTCTGGCCCACAGT
1564	PresenilinI-15	TGCAGGGCTCTCTG
1565	PresenilinI-16	AGTGCAGGGCTCTC
1566	PresenilinI-17	CACTGATCATGATGGC
1567	PresenilinI-18	GACACTGATCATGATGGC
1568	PresenilinI-19	ACAATGACACTGATCATG
1569	PresenilinI-20	GAACCACCAGGAGGAT
1570	PresenilinI-21	GACACAAACAGCCACT
1571	PresenilinI-22	GTGGACCTTTCGGAC
1572	PresenilinI-23	CAACCAGCATACGAAGT
1573	PresenilinI-24	TCCCTCTGGGCTTC
1574	PresenilinI-25	ACTGTCCCTCTGGG .
1575	PresenilinI-26	GACTGTCCCTCTGG
1576	PresenilinI-27	CCTAGATGACTGTCCC
1577	PresenilinI-28	CAGCGAGGATACTGC
1578	PresenilinI-29	CTTCACCAGCGAGGAT
1579	PresenilinI-30	TTTCCTCTGGGTCTTCAC
1580	PresenilinI-31	CTTTCCTCTGGGTCTTC
1581	PresenilinI-32	CTCCCAATCCAAGTTTT
1582	TRADD-1	TTCATCCCGGAGCC
1583	TRADD-2	TTCTTCATCCCGGAGC
1584	TRADD-3	GCTCAGCCAGTTCTTC
1585	TRADD-4	GACAGAGAGGCAC
1586	TRADD-5	CTTCACCTCCGACAG
1587	TRADD-6	GAAAAGTCTGGGCAGG
1588	TRADD-7	GACCCTGGAACAGAAAAG
1589	TRADD-8	CTGACCCTGGAACAG
1590	TRADD-9	ACTACAGGCTGACCCT
1591	TRADD-10	ATTCACTACAGGCTGACC
1592	TRADD-11	CGATTCACTACAGG
1593	TRADD-12	GGCCGATTCACTAC
1594	TRADD-13	CGAACGTCTGTTGGTC
1595	TRADD-14	CGCGAACGTCTGTTG
1596	PKA-1	CTTCTGTTTGTCGAGGAT
1597	PKA-2	TTCACCACCTTCTGTTTG
1598	PKA-3	AGGATGCGCTTTTCATTC
1599	PKA-4	AGCTTGCAGGATGCG
1600	PKA-5	GTTGACAGCTTGCAGGAT

Fig. 5 - 8

1601	PKA-6	GGAACGGAAAGTTGACAG
1602	PKA-7	AACTCGAGTTTGACGAGG
1602	PKA-8	TGTCCTTGAAGGAGAAC
1604	PKA-9	CGTACTCCATGACCATGT
1605	PKA-10	GCACGTACTCCATGAC
1606	PKA-11	GATTCTCCGGCTTCAG
1607	PKA-12	TCAATGAGCAGATTCTCC
1608	PKA-13	GGTCAATGAGCAGATTC
1609	PKA-14	CCCTGCTGGTCAATG
1610	PKA-15	TAGCCCTGCTGGTC
1611	PKA-16	CGCTTGGCGAAACC
1612	PKA-17	CCTTCACGCGCTTG
1613	PKA-18	AAGGTCCAAGTGCG
1614	PKA-19	TGCCGCACAAGGTC
1615	IL-12alpha-1	GGTGAGGACCACCATTT
1616	IL-12alpha-2	GGGTGTCACAGGTG
1617	IL-12alpha-3	ATACCATCTTCTTCAGGG
1618	IL-12alpha-4	GGTGATACCATCTTCTTC
1619	IL-12alpha-5	CCAGGTGATACCATCTTC
1620	IL-12alpha-6	CCTCACTGCTCTGGT
1621	IL-12alpha-7	TAAGACCTCACTGC
1622	IL-12alpha-8	CAGAGCCTAAGACCTC
1623	IL-12alpha-9	CCAGAGCCTAAGACC
1624	IL-12alpha-10	TCTTCCTTTTTGTGAAGC
1625	IL-12alpha-11	GACCAAATTCCATCTTCC
1626	IL-12alpha-12	ATCAGTGGACCAAATTCC
1627	IL-12alpha-13	GGTTCTTTCTGGTCCTTT
1628	IL-12alpha-14	TTTTTGGGTTCTTTCTGG
1629	IL-12alpha-15	GGTCTTATTTTTGGGTTC
1630	IL-12alpha-16	AATGGGCAGACTCTCCT
1631	IL-12alpha-17	TCCACCATGACCTCAATG
1632	IL-12alpha-18	AACGGCATCCACCATG
1633	IL-12alpha-19	GTGAACGGCATCCAC
1634	IL-12alpha-20	ACTTGAGCTTGTGAACGG
1635	IL-12alpha-21	TTCATACTTGAGCTTGTG
1636	IL-12alpha-22	CTGGTGTAGTTTTCATAC
1637	IL-12alpha-23	AGCTGCTGGTGTAGTTTT
1638	IL-12beta-1	AGGAGGACCAGGGT
1639	IL-12beta-2	AGGTGGTCCAGGAG
1640	IL-12beta-3	TTTCTGGCCAAACTGAGG
1641	IL-12beta-4	GGAGGTTTCTGGCC

1642	IL-12beta-5	TCTGGAGTGGCCAC
1643	IL-12beta-6	CTTCTGGAGCATGTTGCT
1644	IL-12beta-7	GCCTTCTGGAGCATG
1645	IL-12beta-8	GTTTGTCTGGCCTTCTG
1646	IL-12beta-9	GAGTTTGTCTGGCCTTCT
1647	IL-12beta-10	CTAGAGTTTGTCTGGCCT
1648	IL-12beta-11	GCAAGGGTAAAATTCTAG
1649	IL-12beta-12	AGTGCAAGGGTAAAATTC
1650	IL-12beta-13	AAACAGGCCTCCACT
1651	IL-12beta-14	CTTGGTTAATTCCAATGG
1652	IL-12beta-15	AGGCAACTCCCATTAGTT
1653	IL-12beta-16	TACTACTAAGGCACAGGG
1654	IL-12beta-17	AATACTACTAAGGCACAG
1655	IL-12beta-18	GTACATCTTCAAGTCTTC
1656	Pg-R	GGAGTGGACATGAT
1657	thr	$\Lambda AGAAGATG\Lambda AGCCTTTG$
1658	ref-fosjun	CCGTCTTACTCTTCTTGG
1659	PIV	CCGATACAATTCCAAGG
1660	PIV	CCTTTTCCTTGAG
1661	PIV	CTGTTGCAAGTACG
1662	bak	CAGAAGCAGAGGC
1663	bak	CCTCAGAAGCAGAGG
1664	bak	CTCCTCAGAAGCAG
1665	bak	ACAGGCTGGTGGCA
1666	bak	CCACTCTCAAACAGGC
1667	bak	ACGGTAGCCGAAGC
1668	bak	GACGGTAGCCGAAGC
1669	bak	GGCCAGACGGTAGC
1670	bak	GTGTAGGGCCAGACGGTA
1671	bak	CCGAAGCCATTTTCAGG
1672	bak	CCCCGAAGCCATTTTC
1673	bak	GGTTGATGTCGTCC
1674	bax	GCTTGAGACACTCGC
1675	bax	CCGGACCCGTCCAT
1676	bclx	GCTTGCTTTACTGC
1677	bclx	GGTTGCTCTGAGAC
1678	bclx	GCCACAGTCATGCC
1679	bmp	CGGGCATGCTGGCG
1680	bmp	GTGAAGTTCAGGATGATC
1681	bmp	CCAGTGCCTCATGG
1682	ICE	CAGTGTTCTCCATGG

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1683	ICE	CTGTACCAGACCGAG
1684	ICE	GCATACTGTTTCAGC
1685	ich	GCCATCAGCTCCTTG
1686	ich	CCACACCATAGATGG
1687	ich	GCTGGAGCAGTTTCC
1688	bcl1	CTCGCTTCTGCTGC
1689	bcl2	ACCGTGGCAAAGCG
1690	mucrep	AGGTGACACCGTGG
1691	AHR	GACTTGATTCCTTCAG
1692	AHR	GGATTTGACTTGATTCC
1693	AHR	GCTGCTGTTCATGG
1694	AHR	CCGTTTCTTTCAGTAGG
1695	CD2	CTTGAAGTAGGAGC
1696	MEK2	CGCTCCTACATGGC
1697	tnf	GATGAGGTACAGGCC
1698	tnf	GTAGATGAGGTACAG
1699	tnf	GAGTAGATGAGGTAC
1700	tnf	CCTGGGAGTAGATG
1701	tnf	GGACCTGGGAGTAG
1702	tnf	ACATGGGTGGAGGG
1703	tnf	GTGCTCATGGTGTC
1704	tnf	CTTTCAGTGCTCATG
1705	tnf	TGCTTTCAGTGCTCA
1706	tnf	GATGATCTGACTGCC
1707	tnf	GTTCGAGAAGATGATC
1708	tnf	GGGTTCGAGAAGATG
1709	tnf	GGTTTGCTACAACATG
1710	tnf	CAGCTTGAGGGTTTG
1711	tnf	TGCCCCTCAGCTTG
1712	TNFR	GACACACTATCTC
1713	IL-18	GCAGCCATCTTTATTC
1714	IL-18	GTTCAGCAGCCATC
1715	IL-18	TGGTTCAGCAGCCA
1716	IL-18	CTACTGGTTCAGCAGC
1717	IL-18	TCTACTGGTTCAGC
1718	IL-18	GCCACAAAGTTGATGC
1719	IL-18	CATTGCCACAAAGTTG
1720	IL-18	GAGAACTTGGTCATTC
1721	IL-18	GGTCAATGAAGAGAAC
1722	IL-18	CGATTTCCTTGGTC
1723	IL-18	CCGATTTCCTTGGTC

1724	IL-18	CAAATAGAGGCCGATTTC
1725	IL-18	CAAATAGAGGCCGA
1726	IL-18	CCTCTAGGCTGGCT
1727	IL-18	CATACCTCTAGGCTG
1728	IL-18	AGCCATACCTCTAG
1729	IL-18	CAGCCATACCTCTAG
1730	IL-18	CACAGAGATAGTTACAG
1731	IL-18	GTCTTCGTTTTGAACAG
1732	IL-18	CTAGTCTTCGTTTTGAAC
1733	IL-18	TAGCTAGTCTTCGTTTTG
1734	IL-18	GAGCCACTGCGCC
1735	IL-18	CGTGAGCCACTGCG
1736	IL-12-Rec	CGTAACGATCACTGG
1737	IL-12-Rec	GCACTCGTAACGATC
1738	IL-12-Rec	GGAGCACTCGTAAC
1739	IL-12-Rec	CATCATCCTGAGGT
1740	IL-12-Rec	CAGTATCATCATCCTG
1741	IL-12-Rec	CTCAGTATCATCC
1742	IL-12-Rec beta2	CTAAAAGTATGTGCCATC
1743	IL-12-Rec beta2	CACATCGCCTCTCT
1744	IL-12-Rec beta2	GCTTCACAGTCACATCGC
1745	IL-12-Rec beta2	GGAAGGCTTCACAGTC
1746	IL-12-Rec beta2	CCTGTGACTTGAGAATTG
1747	IL-12-Rec beta2	GGAAGACCTGTGAC
1748	IL-12-Rec beta2	CTCTGCTCCACATATTTG
1749	IL-12-Rec beta2	CAACGAAGATCTCTG
1750	IL-12-Rec beta2	CAACACCAACGAAG
1751	PKC-beta	GGTCTTCTGTTTGC
1752	CB-1-Rec	CGATGAAGTGGTAGGAAG
1753	TGF-alpha	GGTTGCATGGAAGC
1754	Fascin	GGTCACAAACTTGCC
1755	p300	CTGATTTGGTCCACTAG
1756	CBP	CATGTTAGCACTGTTC
1757	гас-alpha	GGTCTTGATGTACTCC
1758	EBV	CCACCTAAAGAGAGATC
1759	HSPQ	CTTGTACTGCACCATC
1760	CC-CKR1	GCCAGTTAAGAAGATG
1761	CC-CKR4	GAGATCATGATCCATGG
1762	c-CRK	GTAGTGTCCCAATAGTG
1763	c-CRK	CTTCCTCATCATTCCC
1764	CRKL	CACAAGCTTTTCGAC

## **PCT**

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(54) Title: AN ANTISENSE OLIGONUCLEOTIDE PREPARATION METHOD

(57) Abstract

A method for the preparation of an antisense oligonucleotide or derivative thereof comprising the steps of: selecting a target nucleic acid, if necessary elucidating its sequence; generating the antisense oligonucleotide with the proviso that: the oligonucleotide comprises at least 8 residues; the oligonucleotide comprises at maximum twelve elements, which are capable of forming three hydrogen bonds each to cytosine bases; the oligonucleotide does not contain four or more consecutive elements, capable of forming three hydrogen bonds each with four consecutive cytosine bases (CCCC) within the target molecule or alternatively four or more consecutive elements of GGGG; the oligonucleotide does also not contain 2 or more series of three consecutive elements, capable of forming three hydrogen bonds each with three consecutive cytosine bases (CCC) within the target molecule, or alternatively 2 or more series of three consecutive elements of GGG; and the ratio between residues forming two hydrogen bonds per residue (2H-bond-R) with the target molecule and those residues forming three hydrogen bonds per residue (3H-bond-R) with the target molecule, is ruled by the following specifications: 3H-bond-R/3H-bond-R + 2H-bond-R ≥ 0.29; and synthesizing the oligonucleotide thus generated in a per se known manner.

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Inte ional Application No PCT/EP 98/00497

A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 C12N15/11 C07H21/04 A61K31/70 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) C12N C07H A61K IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category ° 1-16 WO 94 25588 A (BIOGNOSTIK GES FUER X BIOMOLEKUL; SCHLINGENSIEPEN GEORG FERDINAN (DE) 10 November 1994 4,6,12 see the whole document, and especially SEQ IDs: 1-56 and 137 for TGF-betal, or SEQ IDs 57 and 136 for TGF-beta2 WO 93 07883 A (ISIS PHARMACEUTICALS INC) 1-4,6-12X 29 April 1993 see page 5, line 20 - page 7 see page 10, line 6 - page 12, line 7 see page 14, line 3 - line 20 6,12 see examples see page 59, line 27 - page 60 see claims Patent family members are listed in annex. Further documents are listed in the continuation of box C. Special categories of cited documents: "I" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such docu-\*O\* document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means.... in the art. \*P\* document published prior to the international filing date but \*&\* document member of the same patent family later than the priority date claimed Date of mailing of the international search report Date of the actual completion of the international search **24**. **0**3. 99 5 November 1998 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, ANDRES S.M. Fax: (+31-70) 340-3016

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Inte 'lonal Application No PCI/EP 98/00497

C.(Continua	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	Polovent to plain No.
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 96 39415 A (ISIS PHARMACEUTICALS INC; CIBA GEIGY (CH); MONIA BRETT P (US); MAR) 12 December 1996 see abstract	1-5, 7-11, 14-17
•	see page 4, line 1 - line 27 see page 9, line 7 - page 14, line 6	
X	YU D ET AL: "HYBRID OLIGONUCLEOTIDES: SYNTHESIS, BIOPHYSICAL PROPERTIES STABILITY STUDIES, AND BIOLOGICAL ACTIVITY" BIOORGANIC & MEDICINAL CHEMISTRY, vol. 4, no. 10, 1996, pages 1685-1692, XP000644792 see the whole document	1-5,7,8, 10,11, 15-17
X	ZHAO Q ET AL: "EFFECT OF DIFFERENT CHEMICALLY MODIFIED OLIGODEOXYNUCLEOTIDES ON IMMUNE STIMULATION" BIOCHEMICAL PHARMACOLOGY, vol. 51, no. 2, 26 January 1996, pages 173-182, XP000610208 see figures 2,3,5,6	1-5, 7-11,17
X	WO 95 00103 A (CHUNG HUN TAEG; IL YANG PHARM CO LTD (KR)) 5 January 1995	1-4, 7-11, 14-17
Α	see pages 6 and 7, SEQ IDs 1,4-8,10-21 see page 7, line 33 - page 10, line 12 see examples 4,5 see claims	13
X	JACHIMCZAK P ET AL: "TRANSFORMING GROWTH FACTOR-BETA-MEDIATED AUTOCRINE GROWTH REGULATION OF GLIOMAS AS DETECTED WITH PHOSPHOROTHIOATE ANTISENSE OLIGONUCLEOTIDES" INTERNATIONAL JOURNAL OF CANCER, vol. 65, no. 3, 26 January 1996, pages 332-337, XP000676566 see the whole document	1-4, 7-11, 13-17
<b>X</b>	HATZFELD J ET AL: "RELEASE OF EARLY HUMAN HEMATOPOIETIC PROGENITORS FROM QUIESCENCE BY ANTISENSE TRANSFORMING GROWTH FACTOR BETA1 OR RB OLIGONUCLEOTIDES" JOURNAL OF EXPERIMENTAL MEDICINE, vol. 174, no. 4, 1 October 1991, pages 925-929, XP002002256 cited in the application see the Rb and p53 antisenses	1-4,7-11
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Int<sup>r</sup> tional Application No PCT/EP 98/00497

		PC1/EP 38/0049/
C.(Continua Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Category	Chadon of doodificity was majoraton, where appropriate, of the relevant paroages	Tielovani to olamii vo.
X	JACHIMCZAK, P. ET AL.: "The effect of transforming growth factor-beta2-specific phosphorothioate anti-sense oligodeoxynucleotides in reversing cellular immunosuppression in malignant glioma"  J.NEUROSURGERY, vol. 78, 1993, pages 944-951, XP002083277 see the whole document	1-4, 7-11,13, 14,17
P,X	FITZPATRICK, D. ET AL.: "Antisense oligonucleotides specific for transforming growth factor beta2 inhibit the growth of malignant mesothelioma both in vitro and in vivo"  CANCER RESEARCH., vol. 57, August 1997, pages 3200-3207, XP002083278 see the whole document	1-5, 7-11,13
A	AGRAWAL S: "Antisense oligonucleotides: towards clinical trials" TRENDS IN BIOTECHNOLOGY, vol. 14, no. 10, October 1996, page 376-387 XP004035728 see table 2 see page 379, left-hand column, line 39 - right-hand column, line 26 see page 383, right-hand column - page 384, right-hand column, paragraph 2	1-17
A	PISETSKY, D. & REICH, C.: "STIMULATION OF IN VITRO PROLIFERATION OF MURINE LYMPHOCYTES BY SYNTHETIC OLIGODEOXYNULEOTIDES" MOLECULAR BIOLOGY REPORT, vol. 18, no. 3, October 1993, pages 217-221, XP000610055 see the whole document	1-17
Α	WO 95 02422 A (WELTMAN JOEL K) 26 January 1995 see the whole document	6,12
Α	WO 96 31600 A (HYBRIDON INC) 10 October 1996 see the whole document	1-17
A	WO 90 10030 A (OLIN CORP) 7 September 1990 see page 4, line 20 - page 7, line 23 see claims	3-6, 10-12
7		

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International application No. PCT/EP 98/00497

### INTERNATIONAL SEARCH REPORT

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Int	ternational Searching Authority found multiple inventions in this international application, as follows:
	see additional sheet
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
	inventions 1. and 39.01 (see continuation-sheet)
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Rema	The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

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INVENTION 1 : Claims 1-17 (all partially)

A method for preparing antisense oligonucleotides and antisenses obtained. Antisense oligonucleotide against the TGF-beta 1 gene and having SEQ ID 41, modified forms thereof, composition containing it and its therapeutic or diagnostic uses.

INVENTIONS 2 to 33: Claims 1-17 (all partially)
As for subject 1, but concerning SEQ IDs 42 to 73 respectively
(invention 2 concerns SEQ ID 42; invention 3, SEQ ID 43 ....;

INVENTION 34: Claims 1-17 (all partially)

invention 33, SEQ ID 73).

Antisense oligonucleotides against the p53 gene, modified forms thereof, composition containing them and their therapeutic or diagnostic uses.

INVENTION 35: Claims 1-17 (all partially)

As for invention 34, but concerning the junB gene.

INVENTION 36: Claims 1-17 (all partially)

As for invention 34, but concerning the junD gene.

INVENTION 37: Claims 1-17 (all partially)

As for invention 34, but concerning the erbB-2 gene.

INVENTION 38: Claims 1-17 (all partially)

As for invention 34, but concerning the c-fos gene.

INVENTION 39.01: Claims 1-17 (all partially)

As for invention 34, but concerning the antisense oligonucleotide against TGF-beta 2 gene and having SEQ ID 519.

INVENTIONS 39.02 to 39.43 : Claims 1-17 (all partially)

As for invention 39.01, but concerning SEQ IDs 520 to 556 and 1273 to 1277 (invention 39.02 concerns SEQ ID 520; invention 39.03, SEQ ID 521.....; invention 39.38, SEQ ID 556; invention 39.39, SEQ ID 1273;...; and invention 39.43, SEQ ID 1277).

INVENTION 40: Claims 1-17 (all partially)

As for invention 34, but concerning the Rb gene.

INVENTION 41: Claims 1-17 (all partially)
As for invention 34, but concerning the relA gene.

INVENTION 42: Claims 1-17 (all partially)
As for invention 34, but concerning the p105/p50 gene.

INVENTION 43: Claims 1-17 (all partially)
As for invention 34, but concerning the NFKB2 gene.

INVENTION 44: Claims 1-17 (all partially)
As for invention 34, but concerning the TANK gene.

INVENTION 45: Claims 1-17 (all partially)
As for invention 34, but concerning the I-kappa B epsilon gene.

INVENTION 46: Claims 1-17 (all partially)
As for invention 34, but concerning the TRAF-6 gene.

INVENTION 47: Claims 1-17 (all partially)
As for invention 34, but concerning the Rank gene.

INVENTION 48: Claims 1-17 (all partially)
As for invention 34, but concerning the IL-5 gene.

INVENTION 49: Claims 1-17 (all partially)
As for invention 34, but concerning the IL-13 gene.

INVENTION 50: Claims 1-17 (all partially)
As for invention 34, but concerning the IL-15 gene.

INVENTION 51: Claims 1-17 (all partially)
As for invention 34, but concerning the I-kappaB(new member) gene.

INVENTION 52: Claims 1-17 (all partially)
As for invention 34, but concerning the Prostaglan.Rec.EP3 gene.

INVENTION 53: Claims 1-17 (all partially)
As for invention 34, but concerning the Presentilin I gene.

INVENTION 54: Claims 1-17 (all partially)
As for invention 34, but concerning the TRADD gene.

INVENTION 55: Claims 1-17 (all partially)
As for invention 34, but concerning the PKA gene.

INVENTION 56: Claims 1-17 (all partially)
As for invention 34, but concerning the IL-12 alpha gene.

- INVENTION 57: Claims 1-17 (all partially)

  As for invention 34, but concerning the IL-12 beta gene.
- INVENTION 5B: Claims 1-17 (all partially)
  As for invention 34, but concerning the Pg-R gene.
- INVENTION 59: Claims 1-17 (all partially)
  As for invention 34, but concerning the thr gene.
- INVENTION 60: Claims 1-17 (all partially)
  As for invention 34, but concerning the ref-fosjun gene.
- INVENTION 61: Claims 1-17 (all partially)
  As for invention 34, but concerning the PIV gene.
- INVENTION 62: Claims 1-17 (all partially)
  As for invention 34, but concerning the bak gene.
- INVENTION 63: Claims 1-17 (all partially)
  As for invention 34, but concerning the bolx gene.
- INVENTION 64: Claims 1-17 (all partially)
  As for invention 34, but concerning the bmp gene.
- INVENTION 65: Claims 1-17 (all partially)
  As for invention 34, but concerning the ICE gene.
- INVENTION 66: Claims 1-17 (all partially)
  As for invention 34, but concerning the ich gene.
- INVENTION 67: Claims 1-17 (all partially)
  As for invention 34, but concerning the bol1 gene.
- INVENTION 68: Claims 1-17 (all partially)
  As for invention 34, but concerning the bcl2 gene.
- INVENTION 69: Claims 1-17 (all partially)

  As for invention 34, but concerning the mucrep gene.
- INVENTION 70: Claims 1-17 (all partially)
  As for invention 34, but concerning the AHR gene.
- INVENTION 71: Claims 1-17 (all partially)
  As for invention 34, but concerning the CD2 gene.
- INVENTION 72: Claims 1-17 (all partially)
  As for invention 34, but concerning the MEK2 gene.

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- INVENTION 73: Claims 1-17 (all partially)
  As for invention 34, but concerning the TNF gene.
- INVENTION 74: Claims 1-17 (all partially)
  As for invention 34, but concerning the TNFR gene.
- INVENTION 75: Claims 1-17 (all partially)
  As for invention 34, but concerning the IL-18 gene.
- INVENTION 76: Claims 1-17 (all partially)
  As for invention 34, but concerning an IL-12-rec gene.
- INVENTION 77: Claims 1-17 (all partially)
  As for invention 34, but concerning the PKC-beta gene.
- INVENTION 78: Claims 1-17 (all partially)
  As for invention 34, but concerning the CB-1-rec gene.
- INVENTION 79: Claims 1-17 (all partially)
  As for invention 34, but concerning the TGF-alpha gene.
- INVENTION 80: Claims 1-17 (all partially)
  As for invention 34, but concerning the Fascin gene.
- INVENTION 81: Claims 1-17 (all partially)
  As for invention 34, but concerning the p300 gene.
- INVENTION 82: Claims 1-17 (all partially)
  As for invention 34, but concerning the CBP gene.
- INVENTION 83: Claims 1-17 (all partially)
  As for invention 34, but concerning the rac-alpha gene.
- INVENTION 84: Claims 1-17 (all partially)
  As for invention 34, but concerning an EBV gene.
- INVENTION 85: Claims 1-17 (all partially)
  As for invention 34, but concerning the HSPQ gene.
- INVENTION 86: Claims 1-17 (all partially)
  As for invention 34, but concerning the CC-CKR1 gene.
- INVENTION 87: Claims 1-17 (all partially)
  As for invention 34, but concerning the CC-CKR4 gene.
- INVENTION 88: Claims 1-17 (all partially)
  As for invention 34, but concerning the c-CRK gene.

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 2	210		
INVENTION 89 : Claims 1-17 (all partially)	RKI dene.		
As for invention 34, but concerning the C	THE GOID.		
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information on patent family members

Inte lional Application No PC1/EP 98/00497

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